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(54) Title: CHIMERIC MOLECULES CONTAINING A MODULE ABLE TO TARGET SPECIFIC CELLS AND A MODULE
REGULATING THE APOPTOGENIC FUNCTION OF THE PERMEABILITY TRANSITION PORE COMPLEX (PTPC)

(57) Abstract: A chimeric polypeptide has the formula: pTox-pTarg, wherein pTox is a viral apoptotic peptide, such as the Vpr peptide of HIV-1 or a fragment of the Vpr peptide of HIV-1 containing the amino acid motif H(F/S)RIG that interacts with mitochondrial inner membrane, adenine nucleotide translocation (ANT) protein of a cell. pTarg is an antibody or an antibody fragment that binds to the outer membrane of the cell. Binding of the chimeric polypeptide to the cell is followed by apoptosis of the cell. A vector encoding a chimeric polypeptide and a recombinant host cell comprising the vector are provided. The chimeric polypeptide is useful for targeting pTox to cells, such as cancer cells.

CHIMERIC MOLECULES CONTAINING A MODULE ABLE TO TARGET
SPECIFIC CELLS AND A MODULE REGULATING THE APOPTOGENIC
FUNCTION OF THE PERMEABILITY TRANSITION PORE COMPLEX (PTPC)

CROSS-REFERENCE TO RELATED APPLICATIONS

The application hereby claims the benefit under 35 U.S.C. § 119(e) of United States provisional application Serial No. 60/265,594, filed February 2, 2001. The entire disclosure of this application is relied upon and incorporated by reference herein.

BACKGROUND OF THE INVENTION

Field of the Invention

The present invention relates generally to cell death regulatory molecules for therapeutic use. More specifically, this invention relates to molecules in which a peptidic or pseudo-peptidic part acting on the permeability transition pore complex (PTPC) is covalently linked to cell-targeting molecules including antibodies, recombinant antibody fragments or homing peptides. The resulting chimeric molecules are polypeptides or peptidomimetic molecules which target the PTPC and/or its major component the adenine nucleotide translocation (ANT) to induce or inhibit cell death (apoptosis). This invention also relates to such chimeric molecules when the PTPC-interacting part is an apoptogenic HIV-1 Vpr-derived peptide (or pseudopeptide) or an ANT-derived peptide (or pseudo-peptide). This invention also relates to nucleic acid sequence construct encoding such chimeric molecule or encoding portions of these chimeric molecules.

Background

It is currently agreed that mitochondria play an important role in controlling life and death of cells (apoptosis; Kroemer and Reed 2000, *Nature Medicine*). It appears both that an increasing number of molecules involved in the transduction of the signal and also many metabolites and certain viral effectors act on mitochondria and influence the permeabilisation of mitochondrial membranes. Using mitochondrial-specific pro-apoptotic agent would seem to be an emerging concept in cancer therapy (Costantini et al 2000, *Journal of the National Cancer Institute*). Similarly, it might be possible to use cytoprotective molecules, thanks to their ability to stabilize mitochondrial membranes, in the treatment of illnesses where there is excessive apoptosis (neurodegenerative diseases, ischemia, AIDS, fulminant hepatitis, etc.).

Mitochondrial membrane permeabilisation (MMP) is a key event of apoptotic cell death associated with the release of caspase activators and caspase-independent death effectors from the intermembrane space, dissipation of the inner transmembrane potential ($\Delta\Psi_m$), as well as a perturbation of oxidative phosphorylation (Green and Reed, 1998; Gross *et al.*, 1999; Kroemer and Reed, 2000; Kroemer *et al.*, 1997; Lemasters *et al.*, 1998; Vander Heiden and Thompson, 1999; Wallace, 1999). Pro- and anti-apoptotic members of the Bcl-2 family regulate inner and outer MMP through interactions with the adenine nucleotide translocation (ANT; in the inner membrane, IM), the voltage-dependent anion channel (VDAC; in the outer membrane, OM), and/or through autonomous channel-forming activities (Desagher *et al.*, 1999; Gross *et al.*, 1999; Kroemer and Reed, 2000; Marzo *et al.*, 1998; Shimizu *et al.*, 1999; Vander Heiden and Thompson, 1999). ANT and VDAC are major components of the permeability transition pore complex (PTPC), a polyprotein structure organized at sites at which the two mitochondrial membranes are apposed (Crompton, 1999; Kroemer and Reed, 2000).

The mitochondrial phase is under the control of Bcl-2 family of oncogenes and anti-oncogenes (for review: 5; 28) involved in more than 50% of cancers (29). All members of Bcl-2 family play an active role in the regulation of apoptosis, some of them being proapoptotic (Bax, Bak, Bcl-X_S, Bad, etc.) and others, being antiapoptotic (Bcl-2, Bcl-X_L, Bcl-w, Mcl-1, etc.) (G. Kroemer, *Nat Med* 3, 614-20 (1997)).

The mitochondrial megachannel is a polyprotein complex formed in the contact site between the inner and the outer mitochondrial membranes that participate in the regulation of mitochondrial membrane permeability. It is composed of a set of proteins including mitochondrion-associated hexokinase (HK), porin (voltage-dependent anion channel or VDAC), adenine nucleotide translocation (ANT), peripheral benzodiazepin receptor (PBR), creatine kinase (CK), and cyclophilin D, as well as Bcl-2 family members. In physiological conditions, PTPC controls the mitochondrial calcium homeostasis via the regulation of its conductance by the mitochondrial pH, the $\Delta\Psi_m$, NAD/NAD(P)H redox equilibrium and matrix protein thiol oxidation. (M. Zoratti, I. Szabo, *Biochim, Biophys Acta* 1241, 139-76 (1995). S. Shimizu, M. Narita, Y. Tsujimoto, *Nature* 399, 483-487 (1999). M. Crompton, *Biochem J* 341, 233-249 (1999). K. Woodfield, A. Ruck, D. Brdiczka, A. P. Halestrap, *Biochem J* 336, 287-90 (1998).

P. Bernardi, K. M. Broekemeier, D. R. Pfeiffer, J Bioenerg Biomembr 26, 509-17 (1994).

F. Ichas, L. Jouaville, J. Mazat, Cell 89, 1145-53 (1997)).

Apoptosis and related forms of controlled cell death are involved in a great number of illness. Excess or insufficiency of cell death processes are involved in auto-immune and neurodegenerative diseases, cancers, ischemia, and pathological infections or diseases such as viral and bacterial infections. Just few examples illustrating the virtually ubiquitous involvement of mitochondria in diseases associated with the abnormal control of cell death will be mentioned here.

In different models of ischemia (heart, liver, kidney or brain), using molecules that are capable of stabilising mitochondrial membranes, such as CsA for example (or its analogous non-immunosuppressor -Me-Val4-CsA) has made it possible to reduce massive apoptosis and its acute consequences at the level of the organ. In addition, VDAC is indispensable for the destruction of neurons of the rat hippocampus after hypoxic reperfusion. In the area of neurodegenerative diseases, a great many observations suggest close links with mitochondrial control of apoptosis (see Kroemer and Reed 2000, Nature Medicine). The neurotoxin -methyl-4-phenylpyridinium induces mitochondrial permeability transition and the exit of cytochrome c. Poisoning by mitochondrial toxins such as nitro-propionic acid or rotenone provokes in primates and rodents a Huntington-disease type of illness.

PTPC is a dynamic protein complex located at the contact site between the two mitochondrial membranes, its opening allowing the free diffusion of solutes < 1500 Da on the inner membrane. Formation of PTPC involves the association of proteins from different compartments, hexokinase (cytosol), porin, also called voltage-dependent anion channel (VDAC, outer membrane), peripheral benzodiazepin receptor (PBR, outer membrane), ANT (inner membrane) and cyclophilin D (matrix). PTPC has been implicated in many examples of apoptosis due to its capacity to integrate multiple pro-apoptotic signal transduction pathways and due to its control by proteins from Bcl-2/Bax family. The Bcl-2 family comprises death inhibitory (Bcl-2-like) and death inducing (Bax-like) members which respectively prevent or facilitate PTPC opening. Bax and Bcl-2 reportedly interact with VDAC and ANT within PTPC. In physiological conditions, ANT is a specific antiporter for ADP and ATP. However, ANT can also form a lethal pore upon interaction with different pro-apoptotic agents. including Ca²⁺,

atractyloside, HIV-1 Vpr-derived peptides and pro-oxidants. Mitochondrial membrane permeabilization may also be regulated by the non-specific VDAC pore modulated by Bcl-2/Bax-like proteins in the outer membrane (12; 16), and/or by changes in the metabolic ATP/ADP gradient between the mitochondrial matrix and the cytoplasm (17).

There is a need in the art for cytoprotective molecules in ischemia, neurodegenerative diseases, fulminant hepatitis and viral infections.

Another application of the chimeric molecule according the invention can be contemplated for the preparation of cosmetics or for preventing early death of plants or vegetables or flowers particularly for preventing the opening of the PTPC.

Conventional chemotherapeutic agents are limited in their therapeutic effectiveness by severe side effects due to their poor selectivity for tumors. The development of monoclonal antibodies (and ScFv) against specific tumor antigens and the identification of homing peptides specific for tumor vascularisation have made it possible to consider enhancing the selectivity of anticancer drugs by a targeted delivery approach. However, such reported attempts using monoclonal antibodies and the anticancer drugs doxorubicin (Trail P.A., et al 1993 Science 261:212), metotrexate (Kanellos J. et al., 1985 J Natl Cancer Inst 75:319), and Vinca alkaloids (Starling J.J. et al., 1991 Cancer Res 41:2965), have been largely unsuccessful. These antibody-drug conjugates were only moderately potent and usually less cytotoxic than the corresponding unconjugated drugs. In fact, antigen-specific cytotoxicity toward cultured tumor cells was rarely demonstrated. *In vivo* therapeutic effects with these conjugates in tumor xenograft animal models were in general observed only when the treatments were commenced before the tumors were well established or when exceedingly large doses (up to 90 mg/kg, drug equivalent dose) were used. It is, therefore, not surprising that in human clinical trials, no significant antitumor effects were observed with these agents (Elias D.J. et al., 1994 Am Respir Crit Care Med 150:1114) (Schneck D. et al., 1990). Indeed, the peak circulating serum concentrations of conjugates were only in the same range as their *in vitro* IC50 value and thus, capable of eliminating at best only about 50% of tumor cells.

These observations led to the conclusion that the previous attempts at delivering therapeutic doses of cytotoxic drugs via monoclonal antibodies have met with little success in clinical trials because of inappropriate choice of drug. One possible (partial-) solution was to

conclude that immunoconjugates must be composed of drugs possessing much higher potency than the clinically used anticancer agents if therapeutic levels of conjugate at the tumor sites are to be achieved in patients. Effectively, such toxins, including maytansinoides, enediyne, or intercalating agents CC1065, were shown to be 100 to 1000-fold more cytotoxic than the chemotherapeutic agents doxorubicin, methotrexate, and Vinca alkaloids (Chari RVJ et al., 1995 *Cancer Res* 55:4079) (Chari RVJ et al., 1992, *Cancer Res* 52:127).

Another approach termed "Adept" was also designed. This antibody-directed enzyme prodrug therapy (Adept) is based upon the use of a monoclonal antibody to target an enzyme at the tumor cell surface, which ultimately is expected to selectively deliver an antitumor drug from a suitable inactive prodrug. In both cases, clinical trials are in progress; however, since today none of them have been introduced in cancer chemotherapy, there is a need for new tools to kill target tumor cells. Bagshawe KD, 1990. Antibody-directed enzyme/prodrug therapy (ADEPT). *Biochem Soc Trans.* 18(5):750-2. Melton RG, Sherwood RF. 1996 Antibody-enzyme conjugates for cancer therapy. *J Natl Cancer Inst.* 88(3-4):153-65. Rihova B. 1997; Targeting of drugs to cell surface receptors. *Crit Rev Biotechnol.* 17(2):149-69. Hudson PJ. 2000. Recombinant antibodies: a novel approach to cancer diagnosis and therapy. *Expert Opin Investig Drugs* 9(6):1231-42.

Recently, the mitochondrion has been proposed as a novel prospective target for chemotherapy-induced apoptosis (1-7). Indeed, four different anti-cancer agents, including the resinoid acid-derivative CD437, lonidamine, betulinic acid, and arsenite, have been shown to induce cancer cell apoptosis by a direct action on mitochondria. The interaction of these anti-cancer agents with mitochondria results in an increase of the permeability of the inner mitochondrial membrane due, at least in part, to the opening of the permeability transition pore complex (PTPC). PTPC opening leads to swelling of the mitochondria matrix, the dissipation of the inner transmembrane potential ($\Delta\Psi_m$), enhanced generation of reactive oxygen species (ROS), and the release of apoptogenic proteins from the intermembrane space to the cytoplasm. Such mitochondrial apoptogenic effectors include the caspase activator cytochrome c, apoptosis inducing factor (AIF), and pro-caspases (2-6). All the signs of apoptosis induced by CD437, lonidamine, betulinic acid, and arsenite are prevented by two agents acting on specific PTPC proteins, namely cyclosporin A (CsA, a cyclophilin D ligand) and bongkrekic acid (BA, a

ligand of the adenine nucleotide translocase (ANT)). It thus appears that PTPC opening is a critical event of apoptosis triggered by these agents.

Mastoparan, a peptide isolated from wasp venom, is the first peptide known to induce mitochondrial membrane permeabilization via a CsA-inhibitable mechanism and to induce apoptosis via a mitochondrial effect when added to intact cells. This peptide has an α -helical structure and possesses some positive charges that are distributed on one side of the helix. A similar peptide (KLAKLAKKKLAKLAK or (KLAKLAK)₂ (K = lysine, L = alanine, and A = leucine) has been found recently to disrupt mitochondrial membranes when it is added to purified mitochondria, although the mechanisms of this effect have not been elucidated.

The vasculature of individual tissues is highly specialized. The endothelium in lymphoid tissues expresses tissue-specific receptors for lymphocyte homing, and recent work utilizing phage homing has revealed an unprecedented degree of specialization in the vasculature of other normal tissues. *In vivo* screening of libraries of phage that displace random peptide sequences on their surfaces has yielded specific homing peptides for a large number of normal tissues. The tissue-specific endothelial molecules to which the phage peptides home may serve as receptors for metastasizing malignant cells. Probing of tumor vasculature has yielded peptides that home to endothelial receptors expressed selectively in angiogenic neovasculature. These receptors, and those specific for the vasculature of individual normal tissues, are likely to be useful in targeting therapies to specific sites. Ruoslahti E, Rajotte D. 2000; An address system in the vasculature of normal tissues and tumors. *Annu Rev Immunol.* 18:813-27.

Ellerby et al. recently have fused the mitochondriotoxic (KLAKLAK)₂ motif to a targeting peptide that interacts with endothelial cells. Such a fusion peptide is internalized and induces mitochondrial membrane permeabilization in angiogenic endothelial cells and kills MDA-MD-435 breast cancer xenografts transplanted into nude mice. Similarly, a recombinant chimeric protein containing interleukin 2 (IL-2) protein fused to Bax selectively binds to and kills IL-2 receptor-bearing cells *in vitro*. Thus, specific cytotoxic agents that target surface receptors, translocate into the cytoplasm, and induce apoptosis via mitochondrial membrane permeabilization might be useful in treating cancer.

There is a need in the art for the selective eradication of transformed cells. One strategy is to target a toxic agent to selected cell types. More particularly, there exists a need in the art for method and reagents for regulating mitochondrial permeabilization and apoptosis.

Summary of the Invention

In order to overcome at least some of the limitations of the prior art, the present invention provides a peptidic or pseudo-peptidic family of polyfunctional molecules containing a cell-targeting part (termed TARG), a PTPC-interacting part (termed TOX/SAVE), and a facultative mitochondrial localisation sequence (MLS). In a preferred embodiment of the invention, the TOX/SAVE portion of the said polyfunctional molecule is a peptide or peptidomimetic molecule which interact directly with the Adenine Nucleotide Translocator (ANT) a central component of the PTPC.

Thus, the present invention includes two categories of targeted cell death regulatory molecules:

- TARG-(MLS)-TOX is a polyfunctional molecule which induces a PTPC-dependent mitochondrial membrane permeabilisation and consequent cell death.
- TARG-(MLS)-SAVE is a polyfunctional molecule which protects cells from mitochondrial membrane permeabilisation and consequently from cell death through interaction with the PTPC and/or ANT.

The invention further provides a vector encoding a chimeric polypeptide of the invention. Also, the invention provides a recombinant host cell comprising a vector of the invention.

Further, the invention provides a cancer cell having a tumor-associated antigen on the surface thereof to which the chimeric polypeptide of the invention is bound via the antibody or antibody fragment of the chimeric polypeptide. The invention also provides methods for detecting cancer cells.

The invention also provides methods for inducing or preventing apoptosis with polypeptides of the invention. The invention provides methods for inducing apoptosis in tumor cells. The invention provides methods for inducing apoptosis in virus infected cells.

The invention further provides hybridomas producing polypeptides of the invention. The invention also provides monoclonal antibodies produced by these hybridomas.

The invention also provides methods for identifying active agents of interest that interact with the PTPC. The invention also provides methods for identifying active agents of interest that interact with ANT peptide. The invention also provides methods for identifying mitochondrial antigens.

The invention also provides methods of treatment or prevention of a pathological infection or disease by administering a polypeptide of the invention to a patient. The invention also provides pharmaceutical compositions comprising a polypeptide of the invention.

Brief Description of the Drawings

Figure 1 shows the nucleotide sequence of vector pACgp67-ScFv461.

Figure 2 shows the nucleotide sequence of vector pACgp67-ScFv350.

Figure 3 shows the nucleotide sequence of Vh and VL, from the clone therap 99B3.

Figure 4 shows the nucleotide sequence of Vh and VL from the clone therap.88E10.

Figure 5 shows the nucleotide sequence of Vh and VL from the clone therap.152C3.

Figure 6, 7, 8, 9, 10, 11 show surface plasmon resonance curves.

Figures 12 and 13 show the strategy for obtaining the ScFv-transfert vector.

Detailed Description of the Invention

It was recently discovered that the proapoptotic HIV-1-encoded protein Vpr induces mitochondrial membrane permeabilization via its physical and functional interaction with the mitochondrial inner membrane protein ANT (adenine nucleotide translocation, also called ADP/ATP carrier). This was shown using a variety of different techniques: surface plasmon resonance, electrophysiology, synthetic proteoliposomes, studies on purified mitochondria (respirometry, electron microscopy, organellofluorometry), as well as microinjection of intact cells. These discoveries are described in detail in U.S. Provisional Application No. 60/231,539 filed September 11, 2000, the entire disclosure of which is relied upon and incorporated by reference herein.

The present invention pertains to novel cytotoxic conjugates based on the association between a peptidic molecule (named pTox) interacting with the mitochondrial permeability transition pore complex (PTPC) and a molecule (named pTarg) able to target cells. The present

invention also pertains to novel cytoprotective conjugates based on the association between a peptidic molecule (named SAVE) interacting with the mitochondrial permeability transition pore complex (PTPC) and a molecule (named pTarg) able to target the cells to rescue. In a specific embodiment of this invention, a cytotoxic conjugate of the invention includes a viral derived pro-apoptotic peptide.

In one embodiment of the invention, the polyfunctional molecule TARG-(MLS)-TOX is a tumor specific molecule that selectively interact with a tumor cell or a specific mammalian cell type, where the polyfunctional molecule is selectively internalised by the mammalian or tumoral cell type, where the polyfunctional molecule interact with the PTPC and/or ANT and exhibits thereto a strong mitochondrio-toxicity leading to apoptosis or any cell death process.

In one embodiment of the invention, the polyfunctional molecule TARG-(MLS)-TOX exhibits a selective toxicity against angiogenic endothelial cells. In another embodiment of the invention, the polyfunctional molecule TARG-(MLS)-TOX exhibits a selective toxicity against tumor cells.

In one embodiment of the invention, the TARG part of the polyfunctional molecule TARG-(MLS)-TOX is an antibody or a recombinant antibody fragment. In another embodiment of the invention, the TARG part of the polyfunctional molecule TARG-(MLS)-TOX is tumor homing peptide (example; CNGRC peptide; lung-homing peptide CGFECVRQCPERC).

In one embodiment of the invention, the TOX part of the polyfunctional molecule TARG-(MLS)-TOX is a peptide or a peptido-mimetic derived from the C-terminal part (amino-acids 52 to 96) of the HIV-1 Vpr protein.

In one embodiment of the invention, the TOX part of the polyfunctional molecule TARG-(MLS)-TOX is a pro-apoptotic Bcl-2 family member such as the Bax or Bid proteins, or a fragment thereof.

In one embodiment of the invention, the TOX part of the polyfunctional molecule TARG-(MLS)-TOX is a D-peptide, is a Ψ -peptide or a retro-inverso peptide chosen among the group of peptidic sequences described in table 1:

Table I:

Name	TOX Peptidic Sequences
Vpr71-82	HFRIGCRHSRIG

Vpr71-82[R73,77,80K]	HFKIGCKHSKIG
Vpr71-96	HFRIGCRHSRIGIIQQRRTRNGASKS
Vpr71-96[R73,77,80K]	HFKIGCKHSKIGIIQQRRTRNGASKS
Vpr52-96	DTWTGVEALIRILQQLLFIHFRIGCRHSRIGIIQQRRTRNGASKS
Vpr52-96[R73,77,80K]	DTWTGVEALIRILQQLLFIHFKIGCKHSKIGIIQQRRTRNGASKS
Vpr52-96[L60,67A]	DTWTGVEAAIRILQQALFIHFRIGCRHSRIGIIQQRRTRNGASKS
Vpr52-82	DTWTGVEALIRILQQLLFIHFRIGCRHSRIG
Vpr52-82[R73,77,80K]	DTWTGVEALIRILQQLLFIHFKIGCKHSKIG
Histatin5 Candida Albicans	DSHARKRHGYKRKFHEKHHSHRGY
Mastoparan Vespuila Lewisii	INLKALAALAKKIL
hNUR77(555-568)	LSRLLGKLPELRTL
hNTR(368-381) neurotrophin receptor	ATLDALLAALRRIQ
Bid(84-100)	RNIARHLAQVGDSMRDR
Bax(57-72)	KKLSECLKRIGDELDs
Bax(72-87)	GQVGRQLAIIGDDINR
HBX(70-78)	ALRFTSARR
DCC(1376-1390)	KTHVKTASLGLAGKA
ANT ₁ (104-116)	DRHKQFWRYFAGN
ANT ₂ (104-116)	DKRTQFWRYFAGN
ANT ₃ (104-116)	DKHTQFWRYFAGN
ANT ₁ (104-116 [A114P])	DRHKQFWRYFPGN
ANT ₂ (104-116)[A114P]	DKRTQFWRYFPGN
ANT ₃ (104-116)[A114P]	DKHTQFWRYFPGN
ANT _{1,2,3} (117-134)	LASGGAAGATSLCFVYPL
ANT ₁ (104-134)	DRHKQFWRYFAGNLASGGAAGATSLCFVYPL
ANT ₂ (104-134)	DKRTQFWRYFAGNLASGGAAGATSLCFVYPL
ANT ₃ (104-134)	DKHTQFWRYFAGNLASGGAAGATSLCFVYPL
ANT ₁ (104-134)[A114P])	DRHKQFWRYFPGNLASGGAAGATSLCFVYPL
ANT ₂ (104-134 [A114P])	DKRTQFWRYFPGNLASGGAAGATSLCFVYPL
ANT ₃ (104-134) [A114P])	DKHTQFWRYFPGNLASGGAAGATSLCFVYPL
Vpr 52-96 [C76S]	DTWTGVEALIRILQQLLFIHFRIGSRHSRIGIIQQRRTRNGASKS
HTLV-1p13II	₁₉ PSLRVWRLCARRLV ₃₂

Bad103-127	NLWAAQRYGRELRRMSDEFVDSFKK
Bax52-76	QDASTKKLSECLKRIGDELDNSMEL

In one embodiment of the invention, the SAVE part of the polyfunctional molecule TARG-(MLS)-SAVE is a L-peptide, a D-peptide or a retro-inverso peptide chosen among the group of peptidic sequences described in table II:

Name	SAVE Peptidic Sequences
ANT ₁ (104-116)	DRHKQFWRYFAGN
ANT ₂ (104-116)	DKRTQFWRYFAGN
ANT ₃ (104-116)	DKHTQFWRYFAGN
ANT _{1,2,3} (117-134)	LASGGAAGATSLCFVYPL
ANT ₁ (104-134)	DRHKQFWRYFAGNLASGGAAGATSLCFVYPL
ANT ₂ (104-134)	DKRTQFWRYFAGNLASGGAAGATSLCFVYPL
ANT ₃ (104-134)	DKHTQFWRYFAGNLASGGAAGATSLCFVYPL

In one embodiment of the invention, the TARG part of the polyfunctional molecule TARG-(MIS)-SAVE is a L-peptide, a D-peptide or a retro-inverso peptide chosen among the group of peptidic sequences described in table III:

ANTENNAPIEDIA third helix (residues 43- 58)	RQIKITFQNRRMKT
HIV-1 Vpr 83-96 transduction domain	IIQQRRTRNGASKS
HIV-1 Tat48-59 transduction domain	GRKKRRQRRRPP
HIV-1 Tat49-57 transduction domain	RKKRRQRRR
pep-1	KETWWETWWTEW

In one embodiment of the invention, the Targ part of the polyfunctionnal molecule TARG-(MLS)-TOX is the decanoic acid $\text{CH}_3(\text{CH}_2)_8\text{CO}-$.

In one embodiment of the invention, the TARG part of the polyfunctional molecule TARG-(MLS)-TOX is an antibody, a recombinant antibody, a recombinant antibody fragment or a ScFv (single chain fragment variable).

In one embodiment of the invention, the TARG part of the polyfunctional molecule TARG-(MLS)-TOX is encoded by the following vector pACgp67-ScFv461 (figure 1).

In one embodiment of the invention, the TARG part of the polyfunctional molecule TARG-(MLS)-TOX is encoded by the following vector pACgp67-ScFv350 (figure 2).

In one embodiment of the invention, the TARG part of the polyfunctional molecule TARG-(MLS)-TOX is a tumor homing peptide as defined by Ellerby et al in PCT/US00/01602.

In one embodiment of the invention, the TARG part of the polyfunctional molecule TARG-(MLS)-TOX/SAVE is a brain or kidney homing peptide as defined by Pasqualini R, Ruoslahti (in *Nature* 1996 Mar 28;380(6572):364-6. Organ targeting in vivo using phage display peptide libraries).

In one embodiment of the invention, pTox is the Vpr peptide of HIV-1 or a fragment thereof. Protein R (Vpr) of human immunodeficiency virus type 1 (HIV-1) is a virion-associated viral gene product with an average length of 96 amino acids, and a molecular weight of approximately 15 kD. Vpr is a highly conserved viral protein among HIV, simian immunodeficiency viruses (SIV). *See* Yuqi Zhao and Robert T. Elder, "Yeast Perspectives on HIV-1 VPR," *Frontiers in Bioscience* 5, d905-916, December 1, 2000.

Vpr has been characterized as an oligomer, and is thought to be divided into three domains on the basis of its structural features: an amino-terminal, negatively charged region that is predicted to form an amphipathic α helix (amino acids 17 to 34); a central hydrophobic domain (amino acids 35 to 75); and a carboxy-terminal, positively charged domain (amino acids 80 to 96). Mutational analysis of Vpr suggests that the nuclear import, virion incorporation, and cell cycle arrest of Vpr are mediated by the distinct functional domains. A structural motif within an amino-terminal helix appears to be important for packaging of Vpr into virions and for maintaining the stability of the protein. A central hydrophobic region, especially the leucine-isoleucine (LR) domain, is reported to be involved in the nuclear localization of Vpr. The cell

cycle arrest function of Vpr was found to be largely located within a carboxy-terminal, positively charged region. *See* Tomoyuki Yamaguchi, Nobumoto Watanabe, Hiromitsu Nakauchi, and Atsushi Koito, "Human Immunodeficiency virus type 1 Vpr Modifies Cell Proliferation via Multiple Pathways," *Microbiol, Immunol.*, 43(5), 437-447, 1999.

The amino acid sequence of human immunodeficiency virus type 1 viral protein R (Vpr) is shown below:

MEQAPEDQGPQREPYNNEWTLELLEELKSEAVRHFPRIWLHNLGQHIYE
TYGDTWAGVEAIIRILQQLLFIHFRIGCRHSRIGVTRQRRARNGASRS.

Vpr and peptides containing conserved H(F/S)RIG repeat motifs can rapidly penetrate human CD4 cells, and cause mitochondrial dysfunction and death by apoptosis. More particularly, recombinant Vpr and C-terminal peptides of Vpr containing the conserved sequence HFRIGCRHSRIG can cause permeabilization of CD4⁺ T lymphocytes, a dramatic reduction of mitochondrial membrane potential, and finally cell death. Vpr and Vpr peptides containing the conserved sequence rapidly penetrate cells, co-localize with the DNA, and cause increased granularity and formation of dense apoptotic bodies. Vpr treated cells undergo apoptosis, and this was confirmed by demonstration of DNA fragmentation. *See* C. Arunagiri, I. Macreadie, D. Hewish and A. Azad, "A C-terminal domain of HIV-1 accessory protein Vpr is involved in penetration, mitochondrial dysfunction and apoptosis of human CD4⁺ lymphocytes," *Apoptosis* 1997; 2: 69-76.

Using a yeast model system, it has been confirmed that there is a cytotoxic activity associated with the C-terminal portion of Vpr, particularly the sequence HFRIGCRHSRIG. Vpr and portions of Vpr containing the sequence HFRIGCRHSRIG can kill a range of mammalian cells including human lymphocytes. *See* I.G. Macreadie, A. Kirkpatrick, P.M. Strike, and A.A. Azad, "Cytotoxic Activities of HIV-1 VPR and Sac1p peptides Bioassayed in Yeast," *Protein and Peptide Letters*, Vol. 4, No. 3, pp. 181-186, 1997.

The C-terminal moiety (Vpr52-96), within an α -helical motif of 12 amino acids (Vpr71-82), contain several critical arginine (R) residues (R73, R77, R80), which are strongly conserved among different pathogenic HIV-1 isolates. L.G. Macreadie, et al., *Proc. Natl. Acad. Sci. USA* 92, 2770-2774 (1995). I.G. Macreadie, et al., *FEBS Lett.* 410, 145-149 (1997). E. Jacotot, et al., *J. Exp. Med.* 191, 33-45 (2000). Thus, the pro-apoptotic portion (pTox) of the chimeric

polypeptide of the invention can contain, for example, the sequence HFRIGCRHSRIG (HIV-1 Vpr71-82), HFKIGCKHSKIG, Vpr 71-96, Vpr 52-96, or a pseudo peptidic variant such as D[HFRIGCRHSRIG].

Other variants of Vpr peptides can also be employed in this invention. Peptide fragments of Vpr encompassing a pair of H(F/S)RIG sequence motifs (residues 71-75 and 78-82 of HIV-1 Vpr) have been shown cause cell membrane permeabilization and death in yeast and mammalian cells. Peptide Vpr⁵⁹⁻⁸⁶ (residues 59-86 of Vpr) forms an α -helix encompassing residues 60-77, with a kink in the vicinity of residue 62. It has been shown that the first of the repeated sequence motifs (HFRIG) participates in a well-defined α -helical domain, whereas the second (HSRIG) lay outside the helical domain and forms a reverse turn followed by a less ordered region. On the other hand, peptides Vpr⁷¹⁻⁸² and Vpr⁷¹⁻⁹⁶, in which the sequence motifs are located at the N-terminus, were largely unstructured under similar conditions, as judged by their C²H chemical shifts. Thus, it has been shown that the HFRIG and HSRIG motifs adopt α -helical and turn structures, respectively, when preceded by a helical structure, but are largely unstructured in isolation. There are implications of these findings for interpretation of the structure-function relationships of synthetic peptides containing these motifs. For example, since the HFRIG and HSRIG sequence motifs adopt helical and turn structures, respectively, when preceded by a helical structure, as in full-length Vpr, but are largely unstructured in isolation, 7-8 residues, sufficient to support at least 1-2 turns of helix, should be included at the N-terminus of Vpr when used as the pTox component of the chimeric polypeptides of the invention to ensure that they are able to adopt the same structure as in the full-length protein. *See* Shenggen Yao, Allan M. Torres, Ahmed A. Azad, Ian G. Macreadie and Raymond S. Norton, "Solution Structure of Peptides from HIV-1 Vpr Protein that Cause Membrane Permeabilization and Growth Arrest," *J. Peptide Sci.* 4: 426-435 (1998). While the Vpr gene codes for a protein of 96-amino-acids, variations have been observed, e.g., Vprs from HIV-1_{HXB2} have 97 and 90-amino-acid residues, respectively. It will be understood that these variants can also be employed in this invention.

For the most effective toxicity, HFRIGCRHSRIG should be surrounded on each side by about eight amino acids from the native sequence. Vpr polypeptides and peptides of greater than 9 amino acids that inhibit or augment Vpr binding, mitochondrial membrane permeabilization, or apoptosis can also be employed in the invention, as well as peptides that are at least 10-20, 20-

30, 30-50, 50-100, and 100-365 amino acids in size. DNA fragments encoding these polypeptides and peptides are encompassed by the invention. Flanking residues should not disrupt the helical structures described above.

The Vpr variants and other viral apoptotic peptides can be assessed for their ability to mediate apoptosis, and thus their suitability for use as pTox in the invention. It is understood that many techniques could be used to assess binding of Vpr or another viral apoptotic peptide to ANT, and that these embodiments in no way limit the scope of the invention. For example, in one embodiment, surface plasmon resonance is used to assess binding of Vpr or another viral apoptotic peptide to ANT. In another embodiment, electrophysiology is used to assess binding of Vpr or another viral apoptotic peptide to ANT. In another embodiment, purified mitochondria are used to assess binding of Vpr or another viral apoptotic peptide to ANT. In another embodiment, synthetic proteoliposomes are used to assess binding of Vpr or another viral apoptotic peptide to ANT. In another embodiment, microinjection of live cells is used to assess binding of Vpr or another viral apoptotic peptide to ANT. These techniques are described in U.S. Provisional Application No. 60/231,539.

In another embodiment, the yeast two-hybrid system developed at SUNY (described in U.S. Patent No. 5,282,173 to Fields et al.; J. Luban and S. Goff, *Curr Opin. Biotechnol.* 6:59-64, 1995; R. Brachmann and J. Boeke, *Curr Opin. Biotechnol.* 8:561-568, 1997; R. Brent and R. Finley, *Ann. Rev. Genet.* 31:663-704, 1997; P. Bartel and S. Fields, *Methods Enzymol.* 254:241-263, 1995) can be used to screen for Vpr-ANT interaction as follows. Vpr, or portions thereof, or another viral apoptotic peptide, responsible for interaction, can be fused to the Gal4 DNA binding domain and introduced, together with an ANT molecule fused to the GAL 4 transcriptional activation domain, into a strain that depends on GAL4 activity for growth on plates lacking histidine. Interaction of the Vpr polypeptide or another viral apoptotic peptide with an ANT molecule allows growth of the yeast containing both molecules and allows screening for the molecules that inhibit or alter this interaction (i.e., by inhibiting or augmenting growth). In an alternative embodiment, a detectable marker (e.g. β -galactosidase) can be used to measure binding in a yeast two-hybrid assay.

Alternatively, the binding properties of Vpr peptide fragments or another viral apoptotic peptide can be determined by analyzing the binding of Vpr peptide fragments or another viral

apoptotic peptide to ANT-expressing cells by FACS analysis. This allows the characterization of the binding of the peptides, and the discrimination of relative abilities of the peptide to bind to ANT. *In vitro* binding assays with Vpr or another viral apoptotic peptide can similarly be used to characterize ANT binding activity.

In another specific embodiment, a cytotoxic conjugate of the invention includes an adenine nucleotide translocation (ANT)-derived pro-apoptotic peptide. The pro-apoptotic portion (pTox) of the conjugate can contain, for example, the sequence DKRTQFWRYFPGN (hANT₂104-116[A114P]) or a pseudo-peptidic variant such as [DKRTQFWRYFPGN].

In another specific embodiment, a cytoprotective conjugate of the invention includes ANT-derived anti-apoptotic peptides. The anti-apoptotic portion (pSave) of the conjugate can contain, for example, the sequence DKRTQFWRYFAGN (hANT₂104-116), the sequence LASGGAAGATSLCFVYPL (ANT 117-134) or a pseudo-peptidic variant such as D[DKRTQFWRYFPGN].

The pTarg component of the chimeric polypeptide of the invention can be an antibody or an antibody fragment. The antibody or antibody fragment can be all or part of a polyclonal or monoclonal antibody. The term "antibodies" is meant to include polyclonal antibodies, monoclonal antibodies, fragments thereof, as well as any recombinantly produced binding partners. Antibodies are defined to be specifically binding if they bind with a K_a or greater than or equal to about 10^7 M⁻¹. Affinities of binding partners or antibodies can be readily determined using conventional techniques, for example those described by Scatchard *et al.*, *Ann. N.Y. Acad. Sci.*, 51:660 (1949).

As used herein, the term "antibody fragment" includes the following:

Fc	A constant region dimer lacking C _H 1
Fab	A light chain dimerized to V _H -C _H 1 resulting from papain cleavage; this is monomeric since papain cuts above the hinge cystines
F(ab)' ₂	A dimer of Fab' resulting from pepsin cleavage below the hinge disulfides; this is bivalent and can precipitate antigen

Fab'	A monomer resulting from mild reduction of F(ab)' ₂ : an Fab with part of the hinge
Fd	The heavy chain portion of Fab (V _H -C _{H1}) obtained following reductive denaturation of Fab
Fv	The variable part of Fab: a V _H -V _L dimer
Fb	The constant part of Fab: a C _H 1-C _L dimer
pFc'	A C _{H3} dimer

Fragments of monoclonal antibodies are of particular interest as small antigen targeting molecules. Antibody fragments are also useful for the assembly of the chimeric polypeptides of the invention designed to carry other pTox agents, such as a therapeutic conjugate. For *in vivo* applications, fragments of antibodies are of interest due to their altered pharmacokinetic behavior, which is useful for cancer therapy with cytotoxic agents, and for their rapid penetration into body tissues, which offer advantages for therapy techniques.

An antibody fragment of particular interest for use in the invention is a minimal Fv fragment with antigen-binding activity. The two chains of the Fv fragment are less stably associated than the Fd and light chain of the Fab fragment with no covalent bond and less non-covalent interaction, but nevertheless functional Fv fragments have been expressed for a number of different antibodies. Two strategies can be employed to stabilize the Fv fragments used in the invention: firstly, mutating a selected residue on each of the V_H and V_L chains to a cysteine to allow formation of a disulphide bond between the two domains; and secondly, the introduction of a peptide linker between the C-terminus of one domain and the N-terminus of the other, such that the Fv is produced as a single polypeptide chain known as a single-chain Fv.

Thus, single-chain Fvs (ScFvs), recombinant V_L and V_H fragments covalently tethered together by a polypeptide link and forming one polypeptide chain, are useful in this invention. For expression of Fv genes, several systems can be effectively used, including myeloma cells, insect, yeast, and *Escherichia coli* cells. Expression in *E. coli* has been a frequently used production method, with both intracellular expression and secretion enabling high yields of ScFv to be made.

The production of ScFv molecules requires the identification of a suitable peptide linker to span the 35-40 Å distance between the C-terminus of one domain and the N-terminus of the other and allow correct folding and assembly of the Fv structure. Several different types of linkers have been used and shown to result in functional ScFv. Polypeptides with the average length of 3-18 amino acids are usually used as links. They can be rich in serine and/or glycine residues, which introduce flexibility, or in charged glutamic acid and/or lysine residues, which improve solubility. Linkers can be selected from searching existing protein structures for protein fragments of the appropriate length and conformation, or by designing them *de novo* based on simple, flexible structures, such as the 15 amino acid sequence (Gly₄Ser)₃.

Active single-chain Fv molecules in both of the two possible orientations, V_H-linker-V_L or V_L-linker-V_H are useful in the invention; however, for some antibodies one particular orientation may be preferable as a free N-terminus of one domain, or C-terminus of the other, may be required to retain the native conformation and thus full antigen binding.

The ScFv may be susceptible to aggregation, with dimers, trimers, and multimers formed. The potential of forming dimers or other multimers with very short linkers, or no linker at all, can be exploited to produce stable pTarg structures. Such an approach can also be used to create pTarg molecules with two different binding specificities by fusing the V_H of an antibody of one specificity to the V_L of another and vice versa.

Fv's stabilized by disulphide linkages can also be employed as the pTarg component of the chimeric polypeptide of the invention. The introduction of a disulphide bond between the V_H and V_L domains to form a disulphide-linked Fv requires the identification of residues in close proximity on each chain, which are unlikely to affect directly the conformation of the binding site when mutated to cysteine, and will be capable of forming a disulphide bond without introducing strain into the structure of the Fv. Sites have been identified in both CDR regions and framework regions, which appear to result in the formation of such disulphide bonds and allow the production of stabilized Fv fragments which retain antigen-binding characteristics.

Due to small size, rapid clearance *in vivo*, stability, and easy engineering, ScFvs employed in this invention have various applications in the treatment of diseases, particularly of cancer. ScFvs can exhibit the same affinity and specificity for antigen as monoclonal antibodies. Dozens of ScFvs with different specificities have been constructed. They are useful for genetic

fusion to the potent toxins (pTox). If the monovalency of ScFv is a disadvantage, constructs with di- or multivalency with increased combining efficiency can be employed.

In a preferred embodiment of the invention, the targeting part (pTarg) of the cytotoxic conjugate is a recombinant portion (ScFv) of a tumor specific antibody, such as the ScFv versions of the M350 and V461 monoclonal antibodies. The hybridoma has been deposited at the CNCM on January 24, 2001, under the Accession Number I-2617.

The pTarg component of the chimeric polypeptide of the invention is preferably a monoclonal antibody or a fragment thereof. Monoclonal antibodies to human cell antigens are preferred. Many tumor-associated antigens are now known and characterized, and antibodies to these allow targeting to different tumor types. Useful tumor-associated antigens are absent on normal tissues and present at high levels on tumor cells, preferably homogeneously on all cells of the tumor. Antigen should also not be shed from the tumor into the blood.

Commonly used tumor-associated antigens and examples of antibodies raised against them are described in the following Table.

Antigen	Tumor type	Representative antibody
Tumor-associated glycoprotein 72 (TAG72), 72 kDa glycoprotein	Pancarcinoma	B72.3, CC49
Carcinoembryonic antigen (CEA), 180 kDa glycoprotein	Pancarcinoma	NP-4, A5B7
Polymorphic epithelial mucin (PEM), >100 kDa glycoprotein	Ovarian, breast, lung	HMFG1
Epithelial membrane antigen (EMA), 40 kDa glycoprotein	Colorectal (and other epithelial tumors)	17-1A
epidermal growth factor receptor (EGFR), 175 kDa glycoprotein	Breast, lung	425
p185 ^{HER2} /c-erb-B2		

Antigen	Tumor type	Representative antibody
(185 kDa glycoprotein)	Breast, lung	4D5
Prostate-specific membrane antigen (PSMA), 100 kDa glycoprotein	Prostate	7E11-C5.3
CD33 67 kDa glycoprotein	Myeloid leukemia	P67.6,M195
CD 20 35 kDa glycoprotein	Lymphoma	C2B8
GD2 ganglioside	Melanoma, neuroblastoma	14-18

An important consideration is the absolute amount of antibody localized to the tumor site. Therefore, the ideal molecule would localize to the tumor in large amounts, delivering a high dose of pTox while clearing rapidly from the circulation and the rest of the body, minimizing non-specific toxicity. Intact antibodies typically circulate for a long period of time and accumulate high levels of activity at the tumor site, whereas antibody fragments clear more rapidly, sparing the dose to normal tissues.

The antibody fragments can also be prepared by phage-display technology. Phage display is a selection technique, according to which an antibody fragment (ScFv) is expressed on the surface of the filamentous phage fd. For this, the coding sequence of the antibody variable genes is fused with the gene that encoded the minor coat phage protein III (g3p) located at the end of the phage particle. The fused antibody fragments are displayed on the virion surface and particles with the fragments can be selected by adsorption on insolubilized antigen (panning). The selected particles are used after elution to reinfect bacterial cells. The repeated rounds of adsorption and infection lead to enrichment. Bacterial proteases can cleave the bond between the g3p protein and antibody fragments, which results in the production of soluble antibody fragments by infected bacterial cells. To release the soluble ScFvs, an excision of the g3p gene is made or an amber stop codon between the antibody gene and the g3p gene is engineered.

Immunoglobins and certain variants thereof are known and many have been prepared in recombinant cell culture. For example, see U.S. Patent 4,745,055; EP 256,654; Faulkner *et al.*, *Nature* 298:286 (1982); EP 120,694; EP 125,023; Morrison, *J. Immun.* 123:793 (1979); Köhler *et al.*, *P.N.A.S. USA* 77:2197 (1980); Raso *et al.*, *Cancer Res.* 41:2073 (1981); Morrison *et al.*, *Ann. Rev. Immunol.* 2:239 (1984); Morrison, *Science* 229:1202 (1985); Morrison *et al.*, *P.N.A.S. USA* 81:6851 (1984); EP 255,694; EP 266,663; and WO 88/03559. Reassorted immunoglobulin chains also are known. See for example U.S. patent 4,444,878; WO 88/03565; and EP 68,763 and references cited therein. DNA encoding immunoglobulin light or heavy chain constant regions is known or readily available from cDNA libraries or is synthesized. See for example, Adams *et al.*, *Biochemistry* 19:2711-2719 (1980); Gough *et al.*, *Biochemistry* 19:2702-2710 (1980); Dolby *et al.*, *P.N.A.S. USA*, 77:6027-6031 (1980); Rice *et al.*, *P.N.A.S. USA* 79:7862-7865 (1982); Falkner *et al.*, *Nature* 298:286-288 (1982); and Morrison *et al.*, *Ann. Rev. Immunol.* 2:239-256 (1984). These materials and techniques can be employed to synthesize the pTarg component of the chimeric polypeptide of the invention.

Polyclonal antibodies employed as the pTarg component of the chimeric polypeptide of the invention can be readily generated from a variety of sources, for example, horses, cows, goats, sheep, dogs, chickens, rabbits, mice, or rats, using procedures that are well known in the art. In general, purified cell surface proteins or glycoproteins or a peptide based on the amino acid sequence of cell surface proteins or glycoproteins that is appropriately conjugated is administered to the host animal typically through parenteral injection. The immunogenicity of cell surface proteins or glycoproteins can be enhanced through the use of an adjuvant, for example, Freund's complete or incomplete adjuvant. Following booster immunizations, small samples of serum are collected and tested for reactivity to cell surface proteins or glycoproteins. Examples of various assays useful for such determination include those described in *Antibodies: A Laboratory Manual*, Harlow and Lane (eds.), Cold Spring Harbor Laboratory Press, 1988; as well as procedures, such as countercurrent immuno-electrophoresis (CIEP), radioimmunoassay, radio-immunoprecipitation, enzyme-linked immunosorbent assays (ELISA), dot blot assays, and sandwich assays. See U.S. Patent Nos. 4,376,110 and 4,486,530.

Monoclonal antibodies employed as the pTarg component can be readily prepared using well known procedures. See, for example, the procedures described in U.S. Patent Nos. RE

32,011, 4,902,614, 4,543,439, and 4,411,993; *Monoclonal Antibodies, Hybridomas: A New Dimension in Biological Analyses*, Plenum Press, Kennett, McKearn, and Bechtol (eds.), 1980. Briefly, the host animals, such as mice, are injected intraperitoneally at least once and preferably at least twice at about 3 week intervals with isolated and purified cell surface proteins or glycoproteins, conjugated cell surface proteins or glycoproteins, optionally in the presence of adjuvant. Mouse sera are then assayed by conventional dot blot technique or antibody capture (ABC) to determine which animal is best to fuse. Approximately two to three weeks later, the mice are given an intravenous boost of cell surface proteins or glycoproteins or conjugated cell surface proteins or glycoproteins. Mice are later sacrificed and spleen cells fused with commercially available myeloma cells, such as Ag8.653 (ATCC), following established protocols. Briefly, the myeloma cells are washed several times in media and fused to mouse spleen cells at a ratio of about three spleen cells to one myeloma cell. The fusing agent can be any suitable agent used in the art, for example, polyethylene glycol (PEG). Fusion is plated out in plates containing media that allows for the selective growth of the fused cells. The fused cells can then be allowed to grow for approximately eight days. Supernatants from resultant hybridomas are collected and added to a plate that is first coated with goat anti-mouse Ig. Following washes, a label, such as ¹²⁵I-labeled cell surface proteins or glycoproteins, is added to each well followed by incubation. Positive wells can be subsequently detected by autoradiography. Positive clones can be grown in bulk culture and supernatants are subsequently purified over a Protein A column (Pharmacia).

The monoclonal antibodies for the pTarg component can be produced using alternative techniques, such as those described by Alting-Mees *et al.*, "Monoclonal Antibody Expression Libraries: A Rapid Alternative to Hybridomas", *Strategies in Molecular Biology* 3:1-9 (1990), which is incorporated herein by reference. Similarly, binding partners can be constructed using recombinant DNA techniques to incorporate the variable regions of a gene that encodes a specific binding antibody. Such a technique is described in Larrick *et al.*, *Biotechnology*, 7:394 (1989).

The monoclonal antibodies and fragments thereof employed as the pTarg component include chimeric antibodies, e.g., humanized versions of murine monoclonal antibodies. Such humanized antibodies may be prepared by known techniques, and offer the advantage of reduced immunogenicity when the antibodies are administered to humans. In one embodiment, the

humanized monoclonal antibody comprises the variable region of a murine antibody (or just the antigen binding site thereof) and a constant region derived from a human antibody.

Alternatively, a humanized antibody fragment may comprise the antigen binding site of a murine monoclonal antibody and a variable region fragment (lacking the antigen-binding site) derived from a human antibody. Procedures for the production of chimeric and further engineered monoclonal antibodies include those described in Riechmann *et al.* (*Nature* 332:323, 1988), Liu *et al.* (*PNAS* 84:3439, 1987), Larrick *et al.* (*Bio/Technology* 7:934, 1989), and Winter and Harris (*TIPS* 14:139, May 1993). Procedures to generate antibodies transgenically can be found in GB 2,272,440, US Patent Nos. 5,569,825 and 5,545,806 and related patents claiming priority therefrom, all of which are incorporated by reference herein.

In a further embodiment of the invention, the targeting part (pTarg) of a cytotoxic chimeric polypeptide is a tumor homing peptide. Such a tumor homing peptide include any homing sequence described by Ellerby *et al.*, in example V, VI, VII, VIII of PCT/US00/01602, the entire disclosure of which is relied upon and incorporated by reference herein.

In preferred embodiments of the invention, the chimeric polypeptide has the sequence CNGRCGG-HFRIGCRHSRIG, or CNGRCGG-D[HFRIGCRHSRIG], or CNGRCGG-Vpr52-96, or CNGRCGG-DKRTQFWYFPGN, or CNGRCGG-D[DKRTQFWYFPGN], or ACDCRGDCFCGG-HFRIGCRHSRIG, or ACDCRGDCFCGG-D[HFRIGCRHSRIG], or ACDCRGDCFCGG-Vpr52-96, or ACDCRGDCFCGG-DKRTQFWYFPGN, or ACDCRGDCFCGG-[DKRTQFWYFPGN], or M350/ScFv-HFRIGCRHSRIG, or M350/ScFv-D[HFRIGCRHSRIG] or M350/ScFv-Vpr52-96, or M350/ScFv-DKRTQFWYFPGN, or or M350/ScFv- D[DKRTQFWYFPGN].

Chimeric polypeptides of the invention can be generated by a variety of conventional techniques. Such techniques include those described in B. Merrifield, *Methods Enzymol.* 289:3-13, 1997; H. Ball and P. Mascagni, *Int. J. Pept. Protein Res.* 48:31-47, 1996; F. Molina *et al.*, *Pept. Res.* 9:151-155, 1996; J. Fox, *Mol. Biotechnol.* 3:249-258, 1995; and P. Lepage *et al.*, *Anal. Biochem.* 213: 40-48, 1993.

Peptides can be synthesized on a multi-channel peptide synthesizer using classical Fmoc-based and pseudopeptide synthesis. In one embodiment of the invention, Vpr52-96, Vpr71-96 and Vpr 71-82 and all the Tox, Save and TARG peptides described in Table I, II, III, are

synthesized by solid phase peptide chemistry. After cleavage from the resin, the peptides are purified and analyzed by reverse-phase HPLC. The purity of the peptides is typically above 98% according to HPLC trace. The integrity of each peptide can be controlled by matrix Assisted Laser Desorption Time of Flight spectrometry. To avoid rapid degradation of the peptides in biological fluids, one or several amide bonds could be advantageously replaced by peptide bond isosters like retro-inverso (NH-CO), methylene amino (CH₂-NH), carba (CH₂-CH₂) or carbaza (CH₂-CH₂-N(R)) bonds.

Alternatively, the chimeric polypeptides of the invention can be prepared by subcloning a DNA sequence encoding a desired peptide sequence into an expression vector for the production of the desired peptide. The DNA sequence encoding the peptide is advantageously fused to a sequence encoding a suitable leader or signal peptide. Alternatively, the DNA fragment may be chemically synthesized using conventional techniques. The DNA fragment can also be produced by restriction endonuclease digestion of a clone of, for example HIV-1, DNA using known restriction enzymes (New England Biolabs 1997 Catalog, Stratagene 1997 Catalog, Promega 1997 Catalog) and isolated by conventional means, such as by agarose gel electrophoresis.

In another embodiment, the well known polymerase chain reaction (PCR) procedure can be employed to isolate and amplify a DNA sequence encoding the desired protein or peptide fragment. Oligonucleotides that define the desired termini of the DNA fragment are employed as 5' and 3' primers. The oligonucleotides can contain recognition sites for restriction endonucleases, to facilitate insertion of the amplified DNA fragment into an expression vector. PCR techniques are described in Saiki *et al.*, Science 239:487 (1988); Recombinant DNA Methodology, Wu *et al.*, eds., Academic Press, Inc., San Diego (1989), p. 189-196; and PCR Protocols: A Guide to Methods and Applications, Innis *et al.*, eds, Academic Press., (1990). It is understood of course that many techniques could be used to prepare polypeptide and DNA fragments, and that this embodiment in no way limits the scope of the invention.

Several methods can be used to link TARG to TOX and TARG to SAVE, depending on the particular chemical characteristics of the molecules. For example, methods of linking haptens to carrier proteins as used routinely in the field of applied immunology. In one embodiment, a premade a PTPC regulatory molecule (TOX or SAVE) can be conjugated to an antibody as antibody fragment (pTarg) using, for example, carbodiimide conjugation.

Carbodiimides comprise a group of compounds that have the general formula R-N=C=N-R, where R and R can be aliphatic or aromatic, and are used for synthesis of peptide bonds. The preparative procedure is simple, relatively fast, and is carried out under mild conditions.

Carbodiimide compounds attack carboxylic groups to change them into reactive sites for free amino groups. Carbodiimide conjugation has been used to conjugate a variety of compounds for the production of antibodies.

The water soluble carbodiimide, 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide (EDC) can be useful for conjugating a PTPC regulatory molecule (TOX or SAVE) to an antibody or antibody fragment molecule. Such conjugation requires the presence of an amino group, which can be provided, for example, by a PTPC regulatory molecule (TOX or SAVE), and a carboxyl group, which can be provided by an antibody or antibody fragment.

In addition to using carbodiimides for the direct formation of peptide bonds, EDC also can be used to prepare active esters, such as N-hydroxysuccinimide (NHS) ester. The NHS ester, which binds only to amino groups, then can be used to induce the formation of an amide bond with the single amino group of the oxorubicin. The use of EDC and NHS in combination is commonly used for conjugation in order to increase yield of conjugate formation.

Other methods for conjugating a PTPC regulatory molecule (TOX or SAVE) to an antibody or antibody fragment also can be used. For example, sodium periodate oxidation followed by reductive alkylation of appropriate reactants can be used, as can glutaraldehyde crosslinking. However, it is recognized that, regardless of which method of producing a chimeric polypeptide of the invention is selected, a determination must be made that an antibody or antibody fragment maintains its targeting ability and that a PTPC regulatory molecule (TOX or SAVE) maintains its activity.

The chimeric polypeptide of the invention may further incorporate a specifically non-cleavable or cleavable linker peptide functionally interposed between the PTPC regulatory molecule (TOX or SAVE) (pTarg) and the antibody or antibody fragment (pTox). Such a linker peptide provides by its inclusion in the chimeric construct, a site within the resulting chimeric polypeptide that may be cleaved in a manner to separate the intact PTPC regulatory molecule (TOX or SAVE) from the intact antibody or antibody fragment. Such a linker peptide may be, for instance, a peptide sensitive to thrombin cleavage, factor X cleavage, or other peptidase

cleavage. Alternatively, where the chimeric polypeptide lacks methionine, the antibody or antibody fragment may be separated by a peptide sensitive to cyanogen bromide treatment. In general, such a linker peptide will describe a site, which is uniquely found within the linker peptide, and is not found at any location in either of the TARG, TOX or SAVE fragment constituting the chimeric polypeptide.

Compositions comprising an effective amount of a chimeric polypeptide of the present invention, in combination with other components, such as a physiologically acceptable diluent, carrier, or excipient, are provided herein. The chimeric polypeptide can be formulated according to known methods used to prepare pharmaceutically useful compositions. They can be combined in admixture, either as the sole active material or with other known materials suitable for a given indication, with pharmaceutically acceptable diluents (e.g., saline, Tris-HCl, acetate, and phosphate buffered solutions), preservatives (e.g., thimerosal, benzyl alcohol, parabens), emulsifiers, solubilizers, adjuvants and/or carriers. Suitable formulations for pharmaceutical compositions include those described in *Remington's Pharmaceutical Sciences*, 16th ed. 1980, Mack Publishing Company, Easton, PA.

In addition, such compositions can be complexed with polyethylene glycol (PEG), metal ions, or incorporated into polymeric compounds such as polyacetic acid, polyglycolic acid, hydrogels, dextran, etc., or incorporated into liposomes, microemulsions, micelles, unilamellar or multilamellar vesicles, erythrocyte ghosts or spheroblasts. Such compositions will influence the physical state, solubility, stability, rate of *in vivo* release, and rate of *in vivo* clearance, and are thus chosen according to the intended application.

The compositions of the invention comprising the chimeric polypeptide can be administered in any suitable manner, e.g., topically, parenterally, or by inhalation. The term "parenteral" includes injection, e.g., by subcutaneous, intravenous, or intramuscular routes, also including localized administration, e.g., at a site of disease or injury. Sustained release from implants is also contemplated. One skilled in the pertinent art will recognize that suitable dosages will vary, depending upon such factors as the nature of the disorder to be treated, the patient's body weight, age, and general condition, and the route of administration. Preliminary doses can be determined according to animal tests, and the scaling of dosages for human administration is performed according to art-accepted practices.

Compositions comprising nucleic acids in physiologically acceptable formulations are also contemplated. DNA may be formulated for injection, for example.

In one of its most general applications, the invention relates to a recombinant vector incorporating a DNA segment having a sequence encoding the chimeric polypeptide of the invention. For the purposes of the invention, the term "chimeric polypeptide" is defined as including any polypeptide where at least a portion of a viral apoptotic peptide is coupled to at least a portion of an antibody or antibody fragment. The coupling can be achieved in a manner that provides for a functional transcribing and translating of the DNA segment and message derived therefrom, respectively.

The vectors of the invention will generally be constructed such that the chimeric polypeptide encoding sequence is positioned adjacent to and under the control of an effective promoter. In certain cases, the promotor will comprise a prokaryotic promoter where the vector is being adapted for expression in a prokaryotic host. In other cases, the promoter will comprise a eukaryotic promoter where the vector is being adapted for expression in a eukaryotic host. In the later cases, the vector will typically further include a polyadenylation signal position 3' of the carboxy-terminal amino acid, and within a transcriptional unit of the encoded chimeric polypeptide. Promoters of particular utility in the vectors of the invention are cytomegalovirus promoters and baculovirus promoters, depending upon the cell used for expression. Regardless of the exact nature of the vector's promoters, the recombinant vectors of the invention will incorporate a DNA segment as defined below.

A recombinant host cell is also claimed herein, which incorporates a vector of the invention. The recombinant host cell may be either a eukaryotic cell or a prokaryotic host cell. Where a eukaryotic cell is used, a Chinese Hamster Ovary (CHO) cell has utility. In another embodiment, when used in combination with a baculovirus promoter, the insect cell lines SF9 or SF21 can be used.

This invention will be described in greater detail in the following Examples.

EXAMPLE 1

Obtaining the murine monoclonal antibody (Ac M350)

Human fetal cells were chosen as a source of immunization. It was the well-known similarities between fetal and tumoral antigens which inspired us to use fetal cells as a source of immunization to produce monoclonal antibodies directed against the epitopes present on tumoral cells. Oncofetal antigens are glycoproteins which are present during intra-uterine life; they disappear at birth and can be re-expressed in pathological situations, particularly in malignant tumors. There are many examples of this antigen community, the best known models being fetoprotein which is associated with 70% of liver tumors, and <<embryo tumor antigens>>, which is often used in human clinical practice and which is a monitoring parameter for patients suffering from cancers of the digestive tract.

A. M350 clone production

These fetal cells were obtained from the sterile removal of the mammary buds of 25-week old female fetuses. Once the buds had been mechanically dissociated into 0.5 mm³ fragments, the cells were resuspended in a Dulbecco medium modified with collagenase and hyaluronidase at 37°C and shaken for between 30 minutes and 4 hours after being monitored under the microscope. As soon as organoids appear, the cells were deposited onto Ficoll, washed, then cultured in a calcium-free DMEM-F12 medium, in hepes, insulin, choleric toxin, cortisol. Once the cells were subcultured once a week. Using this technique the cells duplicated 10 to 20 times giving sufficient cells for immunization purposes.

Balb/c mice were immunized four times, intraperitoneally. The fusion was achieved according the classical technic of Kohler and Milstein. The screening was done with fetal mammary cells, adult mammary cells and breast tumors. Several clones appeared and one, M350 clone, was particularly tested on breast tumors and normal breast tissues. 150 tumor sections were tested: (i.e.) infiltrating intra-canalar and intra-lobular adenocarcimonas, infiltrating lobular adenocarcimonas. Tests were performed using an immunoenzymatic technic with alkaline phosphatase. All the tumors tested positive whereas the normal tissues taken from mammary

samples tested in parallel were negative for weakly positive. Each slide of normal tissue contained lobular type epithelial structures and cavities inside the paleal tissue.

B. Other Hybridomes

Obtaining new murine monoclonal antibodies against associated breast tumor antigens.

In this technology, C57/B16 mice were immunized four times, intraperitonaly, with a mixture of three different breast tumor cell lines (MCF7, MDA, ZR75-1). After fusion and screening the specificity was studied on normal breast tissues and malignant tumors, other tumor samples and peripheral blood cells. The Monoclonal antibodies showing surface tumor labeling were chosen.

EXAMPLE 2

A Cell lines and viruses

The insert cells derived from ovarian tissue of Spodoptera frugiperda (Sf9 insect cells, Vaughn et coll., 1977) and insect cells derived from Trichoplusia ni (High Five insect cells) were maintained at 28°C in TC100 medium supplemented with 5% fetal calf serum and were used for the propagation of recombinant baculoviruses and for the production of recombinant proteins. The recombinant baculoviruses are obtained after co-transfection of insect cells with baculovirus viral DNA (Baculogold, Pharmingen) and recombinant transfer vector DNA.

B. Recombinant transfer vector: pVL-PS-gp671

The recombinant transfer vector pVL-PSgp671 derived from transfer vector pVL1392 (Invitrogen) is used as transfer vector to generate recombinant viruses. It includes from 5' to 3' : the peptide signal sequence of gp67 baculovirus glycoprotein, the sequence coding for a His(6)-Tag, the recognition sequence for the Xa Factor, a polylinker region for subcloning the scFv sequence, a link-sequence: GGC required for the covalent association between cytotoxic peptides and ScFv.

The signal peptide sequence from gp67 was added by insertion of a PCR product of gp67 (obtained by PCR from a commercial pcGP67-B plasmid as a template and the PSgp67-Back and PSgp67-For as primers) at the *Bg*/II site of the pVL1392 plasmid. The sequence coding for the His(6)-Tag sequence and the recognition sequence for the Xa factor were then added by using

insertion of oligonucleotides at the 3' end of the gp67 sequence. By the same way the sequence of the peptide motif required for the covalent association between cytotoxic peptides and ScFv: (-Gly-Gly-Cys) was added at the 3' part of the polylinker (the first G is encoded by the last nucleotide of the XmaI site).

Insertion at BamH1 and Bg1I of overaping primers:

Th1: GAT CCC ATC ATC ACC ACC ACC AC (BamHI-His(6))
 Th2: ATT GAA GGA AGA GAATT CCATG (Factor Xa cleavage -EcoRI-NcoI)
 Th3: GCT GCA GCC CGG GGG ATG TTA AA (Pst1 -XmaI -GGS - STOP- BamHI)
 Th4: CTT CCT TCA ATG TGG TGG TGA TGA TGG (link beween Th1 Th2)
 Th5: GGG CTG CAG CCA TGG GAA TTC T (link between Th2 and Th3)
 Th6: GAT CTT TAA CAT CCC CC (link between Th3 and pVL, -pg67)

C Synthesis of ScFv DNA fragment

VH and VL regions of M350:

Total RNA isolated from M350 hybridome have been used as a template for a reverse transcription using oligo (dT) as primers (Reverse Transcription IBI Fermentas). A PCR realized with those cDNAs and specific primers (mouse Ig-Prime-Kit, Novagen) have led to the selective amplification of VH and VL chains. These regions are then cloned in "blunt" in pST-Blue 1 plasmid and sequenced.

VH and VL regions of other hybridoidines:

Total RNA isolated from selected hybridome was used as a template for a reverse transcription using oligo (dT) (Reverse Transcription IBI Fermentas). A PCR with specific primers (mouse Ig-Prime-Kit, Novagen) led to the selective amplification of VH and VL chains. These products are then cloned in pGEMT (TA cloning System front PROMEGA) vector and sequenced. Three new VH and VL sequences were determined from clone therap.99B3 (**Figure 3**), clone therap.88E10 (**Figure 4**), and therap.152C3 (**Figure 5**).

Obtention of the ScFv-transfer vector:

VH-link-VL chimeric DNA were done by fusion-PCR in two steps (**Figure 12**). The first

step added a link-sequence (Gly-Gly-Gly-Gly-Ser) at the 3' of the VH chain and at the 5' end of the VL chain respectively. The second step was a PCR fusion leading to the chimeric DNA: VH-link-VL. The set of primers used in this second step brings a 5' -EcoRI and a 3'-XmaI sites to VH and VL respectively that will be used for the subcloning of the final product in pVL-PSgp671 vector (**Figure 13**).

D Cotransfection and purification of recombinant baculoviruses

Sf9 cells were cotransfected with viral DNA (BaculoGold ; Pharmingen) and recombinant transfer vector DNA (pVL-PSgp671-ScFv) by the lipofection method (Feloner and Ringold, 1989) (DOTAP; Roche). Screening and purification of recombinant viruses were carried out by the common procedure described by Summers and Smith (Summers and Smith, 1987). The recombinant virus was named BAC-PSgp671-scFv and amplified to constitute a viral stock with an MOI of 10^8 .

E Analysis of recombinant proteins

Infected cells were collected, washed with cold phosphate-buffered saline (PBS) and resuspended in sample reducing buffer (Laemmli, 1970). After boiling (100°C for 5 min), proteins samples were resolved by 12.5% sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) under denaturing conditions (Laemmli, 1970). The apparent molecular weight of the protein was checked by coomassie blue staining or the proteins were transferred onto a nitrocellulose filter (Schleicher and Schuell ; BAS 85, 0.45 μ m) with a semidry blotter apparatus (Ancos). The nitrocellulose membrane was then stained with Ponceau Red (Sigma) and subsequently blocked with a solution of Tris-saline buffer (0.05 M Tris-HCl pH7.4, 0.2 M NaCl) containing 0.05% Tween 20 and 5% non fat milk (TS-sat). ScFv was detected using a mouse monoclonal antibody raised against His(6)-Tag (SIGMA) as primary antibody and a sheep anti-mouse immunoglobulin G (IgG)- horseradish peroxidase conjugate as secondary antibody (1: 3000 Amersham). The immunoreactive bands were visualized by using ECL reagents as described by the manufacturer (Amersham).

F Protein production and purification

To obtain viral stock, Sf9 insect cells cultured in IPL41 medium and 5% FCS are infected in exponential phase with the recombinant baculoviruses at MOI1. After a 7-day incubation period at 28° in IPL41 medium with 5% FCS, the supernatant is harvested by centrifugation at 8000 RPM during 15 min. Then High-five insect cells cultured in Xpress media (Biowhitaker) are infected with recombinant baculovirus in exponential phase at MOI 10, following 1h30 of infection High Five cells were harvested by centrifugation and resuspended in Xpress media without serum. After a 4-day period of incubation at 28°C, the supernatant is harvested by centrifugation at 8000 RPM during 15 min. These supernatants are then concentrated by two rounds of ammonium sulfate precipitation. The precipitate obtained by sedimentation is dialyzed during 12 hours and purified using batch of Ni-NTA agarose beads as described by the manufacturer (Qiagen). After dialysis (2 days, PBS, 4°C) and analysis by Coomassie staining purified proteins were used for the covalent association with cytotoxic peptides.

EXAMPLE 3

Method of coupling ScFv to pTox

The peptide was assembled using Fmoc solid phase peptide synthesis, after the last Fmoc deprotection a propionyloxy succinimide ester was allowed to react, in the presence of diisopropyl ethylamine, with the alpha amino group of the peptide. At the end of the reaction (30 min) the peptide resin was washed with methylene chloride and the peptide was classically cleaved and deprotected under acidic conditions. The activated peptide was then purified by HPLC and its integrity was confirmed by mass spectrometry. The activated peptide was then allowed to react with the ScFv with peptide in a molar ratio of 10:1 (pH7, PBS, glass tube over agitation for 3 hours at room temperature). Then, dialysis was done for 48h against PBS a 4°C. Four Tox peptides were coupled to ScFv using this method:

Tox 11	ScFv-M350-Jac5 (Vpr71-96[C761])
Ctr1ToX11I	ScFv-M350-Jac5M (Vpr71-96[C76S;R73,80A])
Tox 12	ScFv-Vpr52-96[C76S]
Ctr1Tox12	ScFv -Vpr52-96[C76S ; R73A; R80A]

EXAMPLE 4**Examples of Targ-Tox or Targ-Save structures**

All the Tox peptides can have a facultative N-terminal biotin and a facultative C-terminal amide fonction. Tox0 is a Tox peptide which does not necessarily require an association with a Targ. Tox1, Tox2, Tox 5, Tox6, Save1, Save2 and their respective control can posses a facultative gly-gly- (-GG-) linker between the Targ and the Tox/Save motif.

Tox0	Biot-DTWTGVEALIRILQQLLFIHFRIGCRHSRIGIIQQRRTNGASKS
Ctr1Tox0	Biot-DTWTGVEALIRILQQLLFHFAIGCRHSAIGIIQQRRTNGASKS

Tox1	Biot- CNGRC-GG-HFRIGCRHSRIG
Ctr1Tox1	Biot- CNGRC-GG-HFAIGCRHSAIG
Ctr2Tox1	Biot-CNGRC-GG-CNGRC
Ctr3Tox1	Biot-GG-HFRIGCRHSRIG
Ctr4Tox1	Biot-CNGRC-GG-Scramble
Ctr5Tox1	Biot-KETWWETWWTEW-GG-HFRIGCRHSRIG

Tox2	Biot-ACDCRGDCFC-GG-HFRIGCRHSRIG
Ctr1Tox2	Biot- ACDCRGDCFC-GG-HFAIGCRHSAIG

Tox5

Tox5	Biot-CNGRC-GG-DKRTQFWRYFPGN (hANT2m)
Ctr1Tox5	Biot-CNGRC-GG-DKRTQFWRYFAGN (hANT2)
Ctr2Tox5	Biot-CNGRC-GG-DRHKQFWRYFPGN (hANT1m)
Ctr3Tox5	Biot-CNGRC-GG-DKHTQFWRYFPGN (hANT3m)
Ctr4Tox5	Biot-GG-DKRTQFWRYFPGN (hANT2m)
Ctr5Tox5	Biot-GG-DRHKQFWRYFPGN (hANT1m)
Ctr6Tox5	Biot-GG-DKHTQFWRYFPGN (hANT3m)
Ctr7Tox5	Biot-CNGRC-GG-Scramble

Tox6

Tox6	Biot-ACDCRGDCFC-GG-DKRTQFWRYFPGN (hANT2m)
Ctr1Tox6	Biot-ACDCRGDCFC-GG-DKRTQFWRYFAGN (hANT2)
Ctr2Tox6	Biot-ACDCRGDCFC-GG-DRHKQFWRYFPGN (hANT1m)
Ctr3Tox6	Biot-ACDCRGDCFC-GG-DKHTQFWRYFPGN (hANT3m)
Ctr4Tox6	Biot-ACDCRGDCFC-GG
Ctr5Tox6	Biot-ACDCRGDCFC-GG-Scramble

Tox 11

Tox 11	ScFv-M350-Jac5(Vpr71-96[C76])
Ctr1Tox11	ScFv-M350-Jac5M(Vpr71-96[C76;R73,80A])

Save1

Save1	Biot-RKKRRQRRR-DKRTQFWRYFAGN (hANT2)
Ctr1Save1	Biot-RKKRRQRRR-DKRTQFWRYFPGN (hANT2m)
Ctr2Save1	Biot-RKKRRQRRR-DRHKQFWRYFAGN (hANT1)
Ctr3Save1	Biot-RKKRRQRRR-DKHTQFWRYFAGN (hANT3)
Ctr4Save1	Biot-RKKRRQRRR
Ctr5Save1	Biot-RKKRRQRRR-Scramble

Save2

Save 2	Biot-RKKRRQRRR-LASGGAAGATSLCFVYPL (hANT[117-134])
Ctr1Save2	Biot-RKKRRQRRR-GAWSNVLRGMGGAFVLVLY (ANTTM6[271-289])
Ctr2Save2	Biot-RKKRRQRRR-scramble

EXAMPLE 5**Evaluation of mitochondrial and nuclear parameters of Apoptosis in cells (cell lines) and cell-free systems****A. Cells**

MCF-7, MDA-MB231, COS and HeLa cells are cultured in complete culture medium (DMEM supplemented with 2 mM glutamine, 10% FCS, 1 mM Pyruvate, 10 mM Hepes and 100 U/ml penicillin/streptomycin). Jurkat cells expressing CD4 and stably transfected with the human Bcl-2 gene or a Neomycin (Neo) resistance vector [Aillet, *et al.*, 1998 J. Virol. 72:9698-9705] only were kindly provided by N. Israel (Pasteur Institute, Paris). Neo and Bcl-2 U937 cells [Zamzami *et al.*, 1995 J. Exp. Med.], and CEM-C7 cells are cultured in RPMI 1640 Glutamax medium supplemented with 10% FCS, antibiotics, and 0.8 µg/ml G418.

The cell tests that have been implemented determine the pathway (intracellular penetration, then subcellular localization) of the candidates, and the apoptotic status ($\Delta\psi_m$, activation and relocalization of cell death effectors, content in nuclear DNA) of the target cell. In order to determine these parameters it is necessary to use fluorescent probes to label the cells and/or the candidates molecules and to implement the following two analytical procedures : multi-parameter cytofluorimetry and fluorescent microscopy. As far as neuroprotection is

concerned, tests were carried out on primary cultures of cortical neuronal cells from mice embryos. As far as cardioprotection is concerned, tests were carried out on primary cultures of cardiomyocytes from mice embryos.

- Intra-cellular pathway tests: the TARG-TOX ou TARG-SAVE peptides coupled either with biotin (detected using fluorochromes conjugated with streptavidin ; or by ligand-blot after subcellular fractioning) or with FITC (detected by direct observation of living cells, videomicroscopy and image analysis) are added to the cells. It is possible to favor the TOX or SAVE mitochondrial routing by inserting mitochondrial addressing signals (the Apoptosis Inducing Factor or ornithin transcarbamylase, for example). Similarly, the mitochondrial routing is evaluated after modifying sequences and certain lateral chains (phosphorylations, methylations), then replacing the peptides by peptidomimetics.

- Multi-parameter analysis of apoptosis on tumoral and endothelial cell lines, and primary neurons. Fluorescent probes will be used to measure the state of the mitochondrial transmembrane potential (JC1, DioC6, mitoTrackers) and nuclear condensation (Hoescht). Similarly, the post-mitochondrial parameters of apoptosis are evaluated using classical hypoploidy tests and cell surface labeling with annexin V-FITC.

In this type of tests, we evaluate either the cytotoxic potential of the TARG-TOX, i.e. their capacity to kill (via a mitochondrial effect) tumoral ou endothelial cell lines (the best TARG-TOX must also kill over-expressing Bcl-2 cell lines); or the cytoprotective potential of the TARG-SAVE when the neurons are subjected to different apoptogenic treatments.

B. Apoptosis Modulation

PBS-washed cells ($1-5 \times 10^5$ /ml) are incubated with (1 to 5 μM) of pTarg-pTox in complete culture medium supplemented or not with cyclosporin A (CsA; 1 μM), bongkrekic acid (BA; 50 μM), and/or the caspase inhibitors N-benzyloxycarbonyl-Val-Ala-Asp.fluoromethylketone (Z-VAD.fmk; 50 μM ; Bachem Bioscience, Inc.), Boc-Asp-fluoromethylketone (Boc-D.fmk), or N-benzyloxycarbonyl-Phe-Ala-fluoromethylketone (Z-FA.fmk; all used at 100 μM added each 24 h; Enzyme Systems). During exposure to pTarg-pTox, human primary PBLs from healthy donors, purified with Lymphoprep (Pharmacia), are cultured in RPMI 1640 Glutamax medium without any addition of serum. In contrast, PHA blasts (24 h of 1 $\mu\text{g}/\text{ml}$ PHA-P [Wellcome Industries]; 48 h with 100 U/ml human recombinant IL-2 [Boehringer Mannheim]) are cultured with 10% FCS.

C. Cytofluorimetric Determinations of Apoptosis-associated Alterations in Intact Cells

For cytofluorometry, the following fluorochromes are employed: 3,3'-dihexyloxacarbo-cyanine iodide (DiOC(6)3; 40 nM) for mitochondrial transmembrane potential ($\Delta\Psi_m$) quantification, hydroethidine (4 μM) for the determination of superoxide anion generation, and propidium. iodide (PI; 5 μM) for the determination of viability (Zamzami, N., *et al.*, 1995. J. Exp. Med. 182:367-377). The frequency of subdiploid cells is determined by PI (50 $\mu\text{g}/\text{ml}$) staining of ethanol-permeabilized cells treated with 500 $\mu\text{g}/\text{ml}$ RNase (Sigma Chemical Co.; 30 min, room temperature [RT]) in PBS, pH 7.4, supplemented with 5 mM glucose (Nicoletti, I. *et al.*, 1991. J. Immunol. Methods. 139:271-280).

D. Fluorescence staining of live cells and immunofluorescence

For the assessment of mitochondrial and nuclear features of apoptosis, cells cultured on a cover slip are incubated with the $\Delta\Psi_m$ -sensitive dyes chloromethyl-X-rosamine (CMXRos; 50 nM; Molecular Probes, Inc.) or 5,5',6,6'-tetrachloro-1,1',3,3'-tetraethylbenzimidazolylcarbocyanine iodide (JC-1, 2 μM , Molecular Probes), the $\Delta\Psi_m$ -insensitive dye Mitotracker green (1 μM ; Molecular Probes, Inc.), and/or Hoechst 33342 (2 μM , Sigma) for 30 min at 37°C in complete culture medium (Marzo, I. *et al.* 1998. Science. 281:2027-2031).

E. For in situ determinations of pTarg-pTox internalisation

For *in situ* determinations of TARG-(MLS)-TOX/SAVE internalisation, cells are incubated at different times with TARG-(MLS)-TOX/SAVE, and then cells are fixed with 4% paraformaldehyde and 0.19% picric acid in PBS (pH 7.4) for 1 h at RT. Fixed cells are permeabilized with 0.1% SDS in PBS at RT (for 5 min), blocked with 10% FCS, and stained with an mAb specific for hexa-histidine tag (clone HIS-1, IgG2a, SIGMA) revealed by a goat anti-mouse PE conjugate [Southern Biotechnology Associates, Inc.], Hsp60 (mAb H4149 [Sigma Chemical Co.], revealed by a goat anti-mouse IgG1 FITC conjugate), cytochrome c oxidase (COX; mAb 20E8-C12 [Molecular Probes, Inc.], revealed by a goat anti-mouse IgG2a FITC conjugate), or when the Targ is a biotinylated peptide, a streptavidin-PE reagent is added 30 min. followed by detection of the fluorescence intensity by fluorescence (and/or confocal) microscopy.

F. Assessment of mitochondrial parameters *in vitro*

Mitochondria are purified from rat liver, as described (Costantini *et al.*, 1996), and resuspended in 250 mM sucrose + 0.1 mM EGTA + 10 mM -tris[hydroxymethyl]methyl-2-Aminoethanesulfonic acid, pH=7.4). For the induction of PT, mitochondria (0.5 mg protein per ml) are resuspended in PT buffer (200 mM sucrose, 10 mM Tris-MOPS (pH 7.4), 5 mM Tris-succinate, 1 mM Tris-phosphate, 2 μ M rotenone, and 10 μ M EGTA-Tris), and monitored in an F4500 fluorescence spectrometer (Hitachi, Tokyo, Japan) for the 90° light scattering of light (545 nm) to determine large amplitude swelling after addition of 2 mM atractyloside (Atr), 1 μ M cyclosporin A (CsA; Novartis, Basel, Switzerland), 5 μ M CaCl₂, and/or 0.5 to 20 μ M of pTarg-pTox or pTarg-pSave. For the determination of the $\Delta\Psi_m$, mitochondria (0.5 mg protein per ml) are incubated in a buffer supplemented with 1 μ M rhodamine 123 (Molecular Probes, Eugene, OR) and the dequenching of rhodamine fluorescence (excitation 505 nm, emission 525 nm) is measured as described (Shimizu *et al.*, 1998). Supernatants from mitochondria (6800 g for 15min; then 20 000 g for 1 h; 4°C) are frozen at -80°C until determination of apoptogenic activity on isolated nuclei, DEVD-afc cleaving activity, and immunodetection of cytochrome c and AIF. Cytochrome c and AIF are detected by means of a monoclonal antibody (clone 7H8.2C12, Pharmingen) and a polyclonal rabbit anti-serum (Susin *et al.* 1999) respectively.

Swelling of isolated mitochondria

Table F1 :

Tox0, Tox1, Tox5, Tox6 induce permeability transition pore (PTP) opening

Name of molecules 5 μ M	Induction of Mitochondrial swelling (sw) +++ rapid sw ; ++ low sw ; + very low sw ; - no sw t 20 min
-	-
Tox0	+++
Tox1	++
Ctr1Tox1	-
Ctr2Tox1	-
Ctr3Tox1	+
Ctr4Tox1	-
Tox5	++
Tox6	++

Table F2:

Save 1 and Save2 inhibit atractyloside-induced PTP opening

Name of molecules	Mitochondrial swelling (sw) %
-	2
Ca 2+ 100 μ M	100
Atractyloside 600 μ M	110
Save I 5 μ M	2
Save I 5 μ M + Atr 600 μ M	5
Save I 20 μ M	12
Save I 20 μ M + Atr 600 μ M	12
Save II 10 μ M	2
Save II 20 μ M	16
Save II 10 μ M + Atr 600 μ M	16
Save II 20 μ M + Atr 600 μ M	16

G. ANT purification and reconstitution in liposomes

ANT was purified from rat heart mitochondria as previously described (8). After mechanical shearing, mitochondria were suspended in 220 mM mannitol, 70 mM sucrose, 10 mM Hepes, 200 μ M EDTA, 100 mM DTT, 0.5 mg/ml subtilisin, pH7.4, kept 8 min on ice and sedimented twice by differential centrifugations (5 min, 500 \times g, and 10 min, 10,000 \times g). Mitochondrial proteins were solubilized by 6% [v:v] Triton X-100 (Boehringer Mannheim) in 40 mM K₂HPO₄, 40 mM KC1, 2 mM EDTA, pH 6.0, for 6 min at RT and solubilized proteins were recovered by ultracentrifugation (30 min, 24,000 \times g, 4°C). Then, 2 ml of this Triton X-100 extract was applied to a column filled with 1 g of hydroxyapatite (BioGel HTP, BioRad), eluted with previous buffer and diluted [v:v] with 20 mM MES, 200 μ M EDTA, 0.5% Triton X-100, pH6.0. Subsequently, the sample was separated with a Hitrap SP column using a FPLC system (Pharmacia) and a linear NaCl gradient (0-1M). Proteins concentration was determined using microBCA-assay (Pierce, Rockfoll, Illinois). Purified ANT and/or recombinant Bcl-2 were reconstituted in PC/cardiolipin liposomes. Briefly, to prepare liposomes, 45 mg PC and 1 mg cardiolipin were mixed in 1 ml chloroform, and the solvent was evaporated under nitrogen. Dry lipids were resuspended in 1 ml liposome buffer (125 mM sucrose + 10 mM -2-hydroxyethylpiperazine-*N*'-2 ethanesulfonic acid; Hepes, pH 7.4) containing 0.3% n-octyl- β -D-pyranoside and mixed by continuous vortexing for 40 min at RT. ANT (0.1 mg/ml) or recombinant Bcl-2 (0.1 mg/ml) were then mixed with liposomes [v:v] and incubated for 20 min at RT. Proteoliposomes were finally dialysed overnight at 4°C.

H. Pore opening assay

ANT-proteoliposomes are sonicated in the presence of 1 mM 4-MUP and 10 mM KC1 (50W, 22sec, Branson sonifier 250) on ice as previously described (28). Then, liposomes were separated on Sepadex G-25 columns (PD-10, Pharmacia) from unencapsulated products. 25 μ l aliquots of liposomes were diluted to 3 ml in 10 mM Hepes, 125 mM saccharose, pH 7.4, mixed with various concentrations of the proapoptotic inducers and incubated for 1 h at RT. Potential inhibitors of mitochondrial membranes permeabilization such as BA, ATP and ADP, were added to the liposomes 30 min before treatment. After addition of 10 μ l-alkaline phosphatase (5 U/ml, Boehringer Mannheim) diluted in liposomes buffer + 0.5 mM MgCl₂, samples were incubated

for 15 min at 37°C under agitation and the enzymatic conversion of 4-MUP in 4-MU was stopped by addition of 150 µl Stop buffer (10 mM Hepes-NaOH, 200 mM EDTA, pH 10). The 4-MU-dependent fluorescence (360/450 nm) was subsequently quantitated (28) using a Perkin Elmer spectrofluorimeter. Atractyloside, a pro-apoptotic permeability transition inducer, was used in each experiment as a standard to determine the 100% response. The percentage of 4-MUP release induced by Vpr-derived peptides or pTarg-ptox was calculated as following : [(fluorescence of liposomes treated by pTarg-ptox - fluorescence of untreated liposomes) / (fluorescence of liposomes treated by atractyloside - fluorescence of untreated liposomes)] x 100.

ANT pore opening assay:

Table H1 : examples of fuctionnal interaction between ANT and Tox or Save constructs.

Tox0 and Tox6 induce ANT-protéoliposomes permeabilisation. Save1 and Save2 block Atractyloside (Atra) -induced ANT-protéoliposomes permeabilisation

molecules	Permeabilisation of ANT - protéoliposomes
-	-
Atra 50µM	+
Atra 100µM	++
Atra 200µM	+++
Tox0 (Biotin-Vpr52-96) 2µM	+++
Tox6 5µM	++
Biotin-Vpr71-96[C76S] 5µM	++
Save1 5µM	-
Atra 200µM + Save1 5µM	-
Save2 5µM	-
Atra 200µM + Save2 5µM	-

I. Binding assays and western blot

Mouse liver mitochondria were isolated as described (zamzami *et al.*, 2000). For the determination of cytochrome C release, supernatants from pTarg-pTox treated mitochondria (6800 g for 15min; then 20 000 g for 1 h; 4°C) were frozen at -80°C until immunodetection of cytochrome c (mouse monoclonal antibody clone 7H8.2CI2, Pharmingen). For binding assays, purified mitochondria were incubated (250 µg of protein in 100 µl swelling buffer) for 30 min at

RT 5 μ M (binding assay) of pTarg-pTox or biotin-pTarg-pTox. Mitochondria were lysed either after incubation with biotinylated Vpr52-96 (upper panel) or lysed before (lower panel) with 150 μ l of a buffer containing 20 mM Tris/HCl, pH 7.6; 400 mM NaCl, 50 mM KCl, 1mM EDTA, 0.2 mM PMSF, aprotinin (100U/ml), 1% Triton X-100 and 20% glycerol. Such extracts were diluted with 2 volumes of PBS plus 1mM EDTA before the addition of 150 μ l avidin-agarose (ImmunoPure, from Pierce) to capture the biotin-labeled Vpr52-96 complexed with its mitochondrial ligand(s) (2 hours at 4 °C in a roller drum). The avidin-agarose was washed batchwise with PBS (5 x 5 ml; 1000 g, 5 min, 4 °C), resuspended in 100 μ l of 2 fold concentrated Laemmli buffer containing 4% SDS and 5 mM β -mercaptoethanol, incubated 10 min at RT and centrifuged (1000 g, 10 min, 4 °C). Finally, the supernatants were heated at 95 °C for 5 min and analysed by SDS-PAGE (12%), followed by Western blot and immunodetection with a rabbit polyclonal anti-serum against human ANT (kindly provided by Dr. Heide H. Schmid; The Hormel Institute, University of Minnesota, MI; Ref).

J. Flow cytometric analysis of purified mitochondria

Mouse liver mitochondria are isolated as described (zamzami *et al.*, 2000). Purified mitochondria are resuspended in PT buffer (200 mM sucrose, 10 mM Tris-MOPS (pH 7.4), 5 mM Tris-succinate, 1 mM Tris-phosphate, 2 μ M rotenone, and 10 μ M EGTA). Cytofluorometric (FACSVantage, Beckton Dickinson) detection is restricted to mitochondria by gating on the FSC/SSC parameters and on the main peak of the FSC-W parameter. Confirmation *a posteriori* of the validity of these double gating is obtained by labeling of mitochondria with the $\Delta\Psi_m$ -insensitive mitochondrial dye MitoTracker® Green (75 nM; Molecular Probes; green fluorescence). To determine the percentage of mitochondria having a low $\Delta\Psi_m$, the $\Delta\Psi_m$ -sensitive fluorochrome JC-1 (200 nM; 570-595 nm) is added 10 min before CCCP or pTarg-pTox molecules. Percentage of mitochondria having a low $\Delta\Psi_m$, is determined in dot-plot FSC/FL-2 (red fluorescence) windows.

K. Cell-free system of apoptosis

AIF activity in the supernatant of mitochondria is tested on HeLa cell nuclei, as described (Susin *et al.*, 1997b). Briefly, AIF-containing supernatants of mitochondria are added to purified HeLa nuclei (90 min, 37 °C), which are stained with propidium iodide (PI; 10 μ g/ml; Sigma Chemical Co.) and analyzed in an Elite II cytofluorometer (Coulter) to determine the frequency

of hypoploid nuclei. In some experiments isolated mitochondria, cytosols from Jurkat or CEM cells (prepared as described (Susin *et al.*, 1997a)), and/or pTarg-pTox are added to the nuclei. Caspase activity in the mitochondrial supernatant was measured using Ac-DEVD-amido-4-trifluoromethylcoumarin (Bachem Bioscience, Inc.) as fluorogenic substrate.

L. Purification and reconstitution of PTPC in liposomes

PTPC from Wistar rat brains are purified and reconstituted in liposomes following published protocols (Brenner *et al.*, 1998; Marzo *et al.*, 1998b). Briefly, homogenized brains are subjected to the extraction of triton-soluble proteins, adsorption of proteins to a DE52 resin anion exchange column, elution on a KC1 gradient, and incorporation of fractions with maximum hexokinase activity into phosphatidylcholine/cholesterol (5: 1, w:w) vesicles by overnight dialysis. Recombinant human Bcl-2 (1-218) lacking the hydrophobic transmembrane domain (Δ 219-239), produced and purified as described (Schendel *et al.*, 1997) are added during the dialysis step at a dose corresponding to 5% of the total PTPC proteins (approximately 10 ng Bcl-2 per mg lipids). Liposomes recovered from dialysis are ultrasonicated. (120 W) during 7 sec in 5 mM malate and 10 mM KCl, charged on a Sephadex G50 columns (Pharmacia), and eluted with 125 mM sucrose + 10 mM HEPES (pH 7.4). Aliquots (approx. 10^7) of liposomes are incubated during 60 min at RT in 125 mM sucrose + 10 mM HEPES (pH 7.4) in the presence or absence of pTarg-pTox, [52-96]Vpr or atractyloside. Then, liposomes are equilibrated with 3,3'dihexylocarbocyanine iodide (DiOC₆(3), 80 nM, 20-30 min at RT; Molecular Probes), and analyzed in a FACS-Vantage cytofluorometer (Becton Dickinson, San José, CA, USA) for DiOC₆(3) retention, as described (Brenner *et al.*, 1998; Marzo *et al.*, 1998b).

Triplicates of 5×10^4 liposomes are analyzed and results are expressed as % of reduction of DiOC₆(3) fluorescence, considering the reduction obtained with 0.25% SDS (15 min, RT) in PTPC liposomes as 100% value.

Examples of specific peptides and constructs relating to this invention that can be utilized in carrying out the foregoing techniques are shown in Tables I, II, and III, as well as any chimeric molecule that is a combination between TARG and TOX or TARG and SAVE peptides or peptidomimetics.

EXAMPLE 6

Surface plasmon resonance indicates that Tox0, Tox1, Tox5, Tox6, Save1 binds purified ANT but not purified VDAC.

Methodology.

Sensor Chips SA (streptavidin coated sensor chips) were used for immobilisation of the different peptides. Tox1 was immobilised at a density of 0.7 ng/mm², Tox0 at a density of 3.7 ng/mm², Ctr1Tox0 at a density of 1.4 ng/mm², Tox5 at a density of 1 ng/mm², Tox6 at a density of 1 ng/mm², Save1 at a density of 1.3 ng/mm², and the control peptide at a density of 0.8 ng/mm². Association and dissociation kinetics of ANT and VDAC interactions were followed at a rate of 10 μ L/min for 10 minutes (5 minutes association and 5 minutes dissociation). The ligand was regenerated with a 1 minute flux of KSCN 3M. The obtained sensorgrams were analysed by the BiAeval 3.1 software using the method of double references (Myszka D.G. 2000. Kinetic, equilibrium and thermodynamic analysis of macromolecular interactions with BIACORE. *Methods Enzymol.* 323:325-340). From the sensorgrams with the ligands were first subtracted the sensorgrams obtained with the corresponding analyte solvents. A second subtraction was performed with the sensorgrams obtained with the control peptide ligand. The control peptide for the Tox and Save peptides was biot-H19C corresponding to the sequence of the β 2-adrenergic receptor (Lebesgue D., Wallukat G., Mijares A., Granier C., Argibay J., and Hoebeke J. (1998) An agonist-like monoclonal antibody to the human β 2-adrenergic receptor. *Eur.J. Pharmacol.* 348:123-133). The control peptide for Tox0 was Ctr1Tox0.

Results.

Figure 6 shows the interaction between ANT and Vpr for 4 ANT concentrations (6.25 to 50 nM). The sensorgrams were best analysed using the simple Langmuir model with drifting baseline and resulted in a Kd of 0.15 nM with a Rmax of 160 ($\chi^2 = 7.24$). The same analysis was performed for the sensorgrams showing the interaction between ANT and Tox1 (**Figure 7**). Studying the VDAC interaction both with Tox0 and Tox1 at VDAC concentrations which were ten times higher (**Figure 8 and 9**), the sensorgrams showed only extremely low association with

the peptide ligand and the obtained curves could not be analysed by the different Langmuir bindings models.

The other peptides were tested for their interaction with ANT at a concentration of 50 nM (**Figure 10**). Purified ANT recognised Tox5, Tox6, and Save1 with relative affinities of respectively 0.1, 0.7, and 0.01 nM. These value being obtained at only one ANT concentration only give the relative affinity of ANT for the three peptides. Again, the use of 50 nM VDAC to interact with the same peptides did not result in any specific binding as shown in **Figure 11**.

The following references have been cited herein. The entire disclosure of each reference cited herein is relied upon and incorporated by reference herein.

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What is claimed is:

1. Method for inducing or preventing the apoptosis of eukaryotic cells comprising the homing on specific tissue cell population of a chimeric bifunctional molecule able to modulate the activity of permeability transition pore complex (PTPC).
2. A method according to claim 1, wherein said chimeric molecules modulate the activity of the permeability transition pore complex (PTPC) of a specific eukaryotic cell by the regulation of opening or the closing of said pore.
3. A method according to claim 1 or 2, wherein said chimeric molecules comprising at least a first functional molecule and a second functional molecule, wherein said first functional molecule has the function to target specifically a tissue cell population, and the second functional molecule has the function to regulate the apoptosis activity linked to the permeability transition pore complex (PTPC) of said specific cells.
4. A method according to claim 3, wherein said chimeric molecules comprising at least a first functional molecule and a second functional molecule, wherein said first functional molecule has the function to target and to enter specifically in a tissue cell population and the second functional molecule has the function to regulate the apoptosis activity linked to the permeability transition pore complex (PTPC) of said specific cells.
5. A method according to claim 3, wherein said chimeric molecules comprising at least a first functional molecule and a second functional molecule, wherein said first functional molecule has the function to target and to enter specifically in a tissue cell population of interest and the second functional molecule has the function to target specifically and inducing or preventing the death of said cells by apoptosis by the regulation of the

opening or the closing of the permeability transition pore complex (PTPC) of mitochondria or a fragment thereof.

6. A method according to claim 4, wherein said chimeric molecule has the formula:

Targ-Tox,

wherein Tox is a viral or a retroviral apoptotic peptide or a peptidomimetic or a fragment of a protein that interacts with permeability transition pore complex (PTPC) of a specific eukaryotic cell to cause apoptosis of the cell; and Targ is chosen from:

an antibody,

an antibody fragment,

arecombinant antibody fragment,

M350/ScFv,

V461/ScFv,

a homing peptide, and

any peptide chosen in Table III,

wherein said molecule binds and enters the cell specifically.

7. A method according to claim 5, wherein said chimeric molecule has the formula

Targ-Save,

wherein Save is a viral or a retroviral or a cellular antiapoptotic peptide or peptidomimetic or a fragment of protein that interacts with permeability transition pore complex (PTPC) of a specific eukaryotic cell to prevent the apoptosis of the cell with the proviso that when Save peptide is a viral peptide, Save is not vMIA protein of Cytomegalovirus; and

Targ is chosen from:

- an antibody,
- an antibody fragment,
- a recombinant antibody fragment,
- M350/ScFv,
- a homing peptide, and
- any peptide chosen in Table III,

wherein said molecule binds and enters the cell specifically.

8. A method according to anyone of claims 1 to 7, wherein said chimeric molecules comprises a Mitochondrial Localisation Sequence (MLS), which has the function to address specifically the second functional molecule to mitochondrial or intermembrane space-of the mitochondria.
9. A method according to claims 1, 2, 3, 4, 5, 6 and 8, wherein Tox is chosen from the group of peptides of Table I.
10. A method according to claims 1, 2, 3, 4, 5, and 7, wherein Save is chosen from the group of peptides of Table II.
11. A method according to any one of claims 1 to 10, wherein the second functional molecule of said chimeric molecules has the function to interact specifically with ANT of the PTPC of mitochondria also refers to as adenine nucleotide translocator isoforms 1, 2, or 3.
12. A chimeric bifunctional molecule capable to enter specifically in a tissue cell population for induce or prevent death of said cell by apoptosis and comprising at least a first functional molecule covalently linked to a second functional molecule, wherein said first

functional molecule has the function to target and to enter specifically in a tissue cell population of interest and the second functional molecule has the function to target specifically and inducing or preventing the death of said cells by apoptosis by the regulation of the opening or the closing of the permeability transition pore complex (PTPC) of mitochondria or a fragment thereof.

13. A chimeric molecule according to claim 12 which has the formula:

Targ-Tox,

wherein Tox is a viral or a retroviral apoptotic peptide or peptidomimetic or a fragment of a protein that interacts with Permeability Transition Pore Complex (PTPC) of a specific eukaryotic cell to cause apoptosis of the cell; and

Targ is chosen from:

an antibody,

an antibody fragment,

a recombinant antibody fragment,

M350/ScFv,

V461/ScFv,

a homing peptide, and

any peptide of Table III,

wherein said molecule binds and enters the cell specifically.

14. A chimeric molecule according to claim 12 which has of the formula

Targ-Save

Wherein Save is a viral or a retroviral or a cellular antiapoptotic peptide or peptidomimetic or a fragment of protein that interacts with Permeability Transition Pore Complex (PTPC) of a specific eukaryotic cell to prevent apoptosis of the cell, with the proviso that when Save peptide is a viral peptide, Save is not vMIA protein of Cytomegalovirus;

and Targ is chosen from:

an antibody,

an antibody fragment,

a recombinant antibody fragment,

M350/ScFv,

a homing peptide, and

any peptide of Table III,

wherein said molecule binds and enters the cell specifically.

15. A chimeric molecule according to any of claims 12 to 14 comprising a mitochondrial localisation sequence (MLS) which has the function to address specifically the second functional molecule to mitochondrial membranes or intermembrane space.
16. A chimeric molecule according to claims 13 or 15, wherein Tox is chosen from the group of peptides of Table I.
17. A chimeric molecule according to claims 14 and 15, wherein wherein Save is chosen from the group of peptides of Table II.
18. A chimeric molecule according to claims 13, 15 and 16, wherein the Targ and Tox peptides are covalently bonded through a peptide linker comprising 3 to 18 amino acids.

19. A chimeric molecule according to claims 14, 15 and 17, wherein the Targ and Save peptides are covalently bonded through a peptide linker comprising 3 to 18 amino acids.
20. A vector encoding a chimeric molecule as claimed in any one of claims 12 to 19.
21. A hybridoma secreting Targ according to claim 13 or 14 and deposited at the National Collection of Culture and Microorganism (C.N.C.M.) on January 24, 2001, under the accession number n° I 2617.
22. A purified monoclonal antibody encoded by the hybridoma of claim 21.
23. A recombinant host cell comprising a vector as claimed in claim 20.
24. A cancer cell having a tumor associated antigen on the surface thereof to which is bound the chimeric molecule as claimed in any one of claims 12 to 19.
25. A method of determining the presence of a cancer cell having a tumor-associated antigen on the surface thereof in a biological sample comprising :
 - a) contacting a biological sample of interest with a chimeric peptide molecule according to claims 12 to 19 under conditions to permit the binding between the chimeric peptide according to the invention and the antigen on the surface of the cancer cell,
 - b) detecting the binding by usual technique; and
 - c) optionally quantifying the binding detected in step b).
26. A method for inducing death by apoptosis in a tumoral or viral infected cell having a tumor-associated antigen on surface thereof in a biological sample comprising: contacting a biological sample of interest with a chimeric peptide molecule according to claims 16 or 17 under conditions to permit the binding between the chimeric peptide

according to the invention and the antigen on the surface of the cancer cell and for a time sufficient to allow the entry inside the cell and death cell by apoptosis or viral infected cells.

27. A method for prevent cell death by mitochondrial apoptosis comprising contacting a biological sample of interest with a chimeric molecule, **molecule** according to claims 17 or 19 under conditions to permit the binding between the chimeric molecule according to the invention and the cell of interest and for a time sufficient to allow the entry inside cell of interest and prevent the cell death by apoptosis.
28. A method for prevent cell death according to claim 27, wherein the cells of interest are choosen among the following cell populations: neurons, cardiocytes, and hepatocytes.
29. A method for identifying an active agent of interest that interacts with the activity of the permeability transition pore complex (PTPC) comprising
 - a) contacting a biological sample containing cells with permeability transition pore complex (PTPC) with a chimeric peptide according to claims 12 to 19 in the presence of a candidate agent; and
 - b) comparing the binding of the chimeric peptide with the permeability transition pore complex (PTPC) in absence of said agent.
 - c) optionally, testing the activity of said selected agent on a preparation of a cellular extract comprising subcellular elements with the permeability transition pore complex (PTPC).

30. A method for identifying an active agent of interest that interacts with ANT peptide of permeability transition pore complex (PTPC) comprising:
 - d) contacting a biological sample containing cells with ANT peptide of permeability transition pore complex (PTPC) with a chimeric peptide according to claims 12 to 19 in the presence of a candidate agent; and
 - e) comparing the binding of the chimeric peptide with the ANT peptide of the permeability transition pore complex (PTPC) in absence of said agent.
 - f) optionally, testing the activity of said selected agent on a preparation of a cellular extract comprising subcellular elements with the ANT peptide of the permeability transition pore complex (PTPC).
31. A method of identification of mitochondrial antigen, said antigen having the capacity to interact with a macromolecule or a molecule or a peptide carrying the characteristic of Tox according to claims 13 or 16.
32. A method of identification of mitochondrial antigen, said antigen having the capacity to interact with a macromolecule or a molecule or a peptide carrying the characteristic of save according to claims 14 or 17.
33. A method of treatment or of prevention of a pathological infection or disease comprising the administration to a patient of the pharmaceutical composition containing at least a chimeric molecule according to any of claims 12 to 19.
34. A pharmaceutical composition comprising at least a chimeric molecule according to claims any of 12 to 19.

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pACgp67-ScFv461 -> 1-phase Translation

DNA sequence 10517 b.p. AAGCTTAACTCG ... AGGGCACTGGC linear

Fig. 1A

1/1 AAG CTT TAC TCG TAA AGC GAG TTG AAG GAT lys leu tyt ser OCH ser glu leu lys asp 91/31 TCG CAC AAC TAT TTA CAA TCC GGC CAA GTT trp his asn tyt leu glu cys gly glu val 181/61 GTG ACT ACG GAA GAT GTG TGG ACC GCA val thr ser glu glu asp val trp thr ala 271/91 AAA TAT TAT GAT GGT GTG CAT TTT TTG CGG lys tyt tyt asp gly val his phe leu arg 361/121 GTT CAA GAA TTT ATT GAC ACG GIA AAA GAA val glu glu phe ile asp thr val lys glu 451/151 ACC CGT TAC ATG GTG TGC AGA TAT TTA ATG CAA CCT CGT ATT CGG CGS CAG GAA ACC ATA thr gly tyt met val cys arg tyt leu met 511/181 AAA ATT GAA AGA CAA AAT TAC GTT CAA GAT lys ile glu arg glu asn tyt val gln asp 631/211 TGT TCC CCT TCT TCC ACC GAA CCA AAA CTA cys cys ala ser ser glu pro lys leu 721/241 GAT TAG CGC GGA TAT TCT CCA CGA CAA TGT ACC GAA COT TGA TGT TAC CTT TAT GCT TTT 811/271 TAG CGG TAA TAC ACC TAG TGC CGT CGG CGG TCA CCG ACA ACA CCG GAT GTT TGC GCT TGT CGG CGG GGT ATT GAA CGG CGC GAT CGA ACA AAT AMB pro OCH thr AMB cys arg arg ala ser 901/301 CCA CCA CTT TGG CAA CTA AAT CGG TGA CCT pro pro leu trp gln leu asn arg OPA pro 991/331 ACT CTT GTT TTT TAA CAA GTT CCT CGG TTT TGT CCT CCA CCA CGG CTT CCA CGG CCT TGT 1081/361 ACT CTT GTT TTT TAA CAA GTT CCT CGG TTT TGT CCT CCA CCA CGG CTT CCA CGG CCT TGT 1171/391 TCA CGA ACT GTT TGC TCT CCT CCT CGG CCT ser pro thr val cys ser pro pro pro val 1251/421 CGC ATT CTT GIA ATT CTA CGG CGT AAG GCA ATT TGG ACT TCA TAA TCA CCT GAA TCA CGG 1351/451 CGA ATT ACA CGG CGT CGG CGG TTT TCA CGA CGC TGT TAG AGG TAG CGC CGG CAT TTT GCA 1531/511 GCT CTA GCA CGT ACC GCA CGT TGA AGG TAT CCT CTC CAA ATT TAA ATT CTC CAA TTT TAA CGC GAG CCA CCT TGT TGA TAC ACC TGT GTC GAT 1621/541 TTT GCA ACA ACT ATT GTT TTT TAA CGG AAA CTA AAC TTA TGC TGG TAA CGA ATA ATT AAA TAT 1711/571 GTC GTT ATT AAC GCA GAC CGC CGG CGT CTC CGC CGA ACC CGC TAA AAC GTG TTG CGC GTT 1801/601 TAC AGT TTT GAT TTG CAT ATT AAC CGC GAT TTT TTA ATT AAC CCT CGG CGC AAG GCA CGT CGG CCT 1891/631 CTC GCT CGA CCT CCA GCA CGT CGT TGA CGC CCT CCT CGG TOT CGC CGA ACA CGT CGA CGG CGT CGT CGA CGG TGA CGA CGG CGC CGC ACC 1981/661 CGA CGC ACA AGT ATT TGT ACA CGG AAT GAT CCT CGG CGC AAG GCA CGT CGG CCT CGA ATT GGC 1991/671 AGT TGG GTT GTT TGC GCA TAT CTA TCG CCT CGT CGG CGA TGT AGC TCC GAA CGT TGA TTT TCA CGA AAG CGG AAA TAA AAC TAA ATT CAT TGC GAT 2151/721 TAG TGG GAT TAA AAC GTT GIA CAT CCT CGC TTT TAA TCA TCC CGT CGA TTA ATT CGC CGA ATT GAC TCA ACT GAT CAA ACT GTG GAA TAA 2251/751 AAT ATT TTA TGT ATT GCA CCT CGA CGG CCT CGG CGC AAG GCA CGT CGG CCT CGA ATT GGC 2341/781 AAT ATT TTA TGT ATT GCA CCT CGA CGG CCT CGG CGC AAG GCA CGT CGG CCT CGA ATT GGC 2372/791 31/11 CAT ATT TAG TTG CGT TTA TGA GAT AAG ATT his ile AMB leu arg leu OPA asp lys ile 211/41 GAA CGC ACG TGT AAA ATG TTT CGC CGG CGT lys ser thr cys lys met phe pro ala arg 151/51 151/51 211/71 241/81 301/101 331/111 391/131 421/141 511/171 571/191 601/201 751/251 781/251 841/281 871/291 931/311 961/321 1021/341 1051/351 1111/371 1141/381 1201/401 1231/411 1291/431 1321/441 1381/461 1411/471 1551/521 1591/531 1591/531 1591/551 1681/551 1741/561 1771/591 1921/561 1951/651 2011/671 2041/681 2101/701 2131/711 2281/761 2311/771 2372/791 2401/801 6/21 GAA CGC ACG TGT AAA ATG TTT CGC CGG CGT 151/51 151/51 211/71 241/81 301/101 331/111 391/131 421/141 511/171 571/191 601/201 751/251 781/251 841/281 871/291 931/311 961/321 1021/341 1051/351 1111/371 1141/381 1201/401 1231/411 1291/431 1321/441 1381/461 1411/471 1551/521 1591/531 1591/531 1591/551 1681/551 1741/561 1771/591 1921/561 1951/651 2011/671 2041/681 2101/701 2131/711 2281/761 2311/771 2372/791 2401/801

pACgp67-ScFv461 -> 1-phase Translation

Fig 1B

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pACgp67-ScFv461 -> 1-phase Translation

Fig. 1C

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5041/1481 STOP
 tgc taa aga tct gat cct TAC CTG GGA CCC GGC AAG AAC CAA AAA CTC ACT CTC TTC AAG GAA ATC CGT ATC GTT AAA CCC GAC ACG AAC
 cys CCA arg ser asp pro phe leu gly pro gly lys ASN glu lys leu Thr leu phe lys glu ile arg ASN val lys Pro asp Thr Met
 5311/1711 5161/1721 5101/1701
 RAG CTT GTC GTT GGA TGG AAA GGA AAA GAG TTC TAC AGC GAA ACT TGG ACT CCC TTC ATG GAA GAC AGC TTC CCC ACT GTT AAC GAC CAA
 lys leu val val cly trp lys gly lys glu Thr Tyr ASN Glu Thr Arg Phe Met Ser Glu Asp Ser Phe Pro Ile Val ASN Val Asp Glu
 5221/1741 5251/1751 5191/1731
 GAA CTG ATG GAT GTT TTC CTT GTC AAC ATG COT CCC ACT AGA CCC AAC COT TGT TAC AAA TAC CGT GCA CAA CAC GCT CTG CGT TCC
 glu val met asp val phe leu val val ASN Met Arg Pro Thr Arg Pro ASN Arg Cys Tyr Lys Phe Leu Ala Glu His Ala Leu Arg Cys
 5311/1771 5341/1781 5371/1791
 GAC CCC GAC TAT GTC CCT CAT GAC GTG ATT AGG ATC GTC GAG CCT TCA TGG GTC GGC AGC AAC AAC GAG TAC CGC ATC AGC CTG OCT AAC
 asp pro asp Tyr Val Pro His Asp Val Ile Arg Ile Val Glu Pro Ser Trp Val Gly Ser ASN ASN Glu Tyr Arg Ile Ser Leu Ala Lys
 5401/1801 5431/1811 5461/1821
 AAG GGC GGC GGC TGC CCA ATA ATG AAC CTT CAC TCT GAG EAC ACC AAC TCG TTC GAA CAG TTC ATC GAT CGT CTC ATC TGG GAG AAC TTC
 lys gly gly gly Cys Pro Ile Met ASN Leu His Ser Glu Tyr Thr ASN Ser Phe Glu Glu His Glu His His Leu Arg Cys
 5491/1831 5521/1841 5551/1851
 TAC AAG CCC ATC CTT TAC ATC GGT ACT GAC TCT GCT GAA GAS GAG GAA ATT CTC CTT GAA GTC TCC CTG GTG TTC AAA GTC AAG GAG TTC
 Tyr Lys Pro Ile Val Tyr Ile Gly Thr Asp Ser Ala Glu Glu Glu Ile Leu Glu Val Ser Ile Val Phe Lys Val Lys Glu Phe
 5581/1851 5611/1871 5641/1881
 GCA CCA GAC CCA CCT CTG TTC ACT GGT CCG GCG TAT TAA AAC ACG ATA CAT TGT TAT TAG TAC ATT TAT TAA GCG CTA GAT TCT CTG CTC
 Ala Pro Asp Ala Pro Leu Phe Thr Gly Pro Ala Tyr OCH ASN Thr Ile His Cys Tyr ASN Tyr Ile Tyr OCH Ala Leu Asp Ser Val Arg
 5671/1891 5701/1901 5731/1911
 TGT TGA TTT ACA GAC AAT TGT TGT ACC TAT TTT ATT ATT TCA TAA ATT TTA TAA TCT TTA GGC TGG TAT GTT AGA GGC AAA ATC AAA TGA
 Cys OPA Phe Thr Asp ASN Cys Thr Tyr Phe ASN ASN Ser Leu ASN Leu ASN OCH Ser Leu Glu Trp Tyr Val Arg Ala Lys Ile Lys OPA
 5761/1921 5791/1931 5821/1941
 TTT TCA GCG TCT TTA TAT CTG AAT TTA ATT ATT AAA TCC TCA ATA GAT TIG TAA ATT AGC TTT CGA TTA GTT TCA AAC AAG GGT TGT TTT
 Phe Ser Ala Ser Leu Tyr Leu ASN Leu ASN Ile Lys Ser Ser Ile Asp Leu OCH ASN Arg Phe Arg Leu Val Ser ASN Lys Gly Cys Phe
 5851/1951 5881/1961 5911/1971
 TCC GAA CCTG ATG CCT GGA CTA TCT ATT GGA TTT TCG CTC AAC GCC ACA AAA CTT GCC AAA Ser Glu Pro Met Ala Gly Leu Ser ASN Gly
 Phe Ser Leu ASN Gly Phe Ser Leu ASN Ala Thr Lys Leu Ala Iys 5971/1981 5971/1991 6001/2001
 TTC GTT TGT GTT TTG TTT ATT AAA GGT TCG ACG TCG TTC AAA ATA TTA TGG CCT TTT GTC ATT TTT GTC ATT TCT TCA TCA CTG TCG TTA GTC
 Phe Val Cys Val Leu Phe Cys ASN Lys Gly Ser Thr Ser Phe Lys Ile Leu Cys Ala Phe Val Phe Leu Ser Ser Leu Val Val Tyr
 6031/2011 6061/2021 6091/2031
 ATT TGA CTC GAC GTC AAC AGC TTA ATT AAA GCT TGG ACA ATT TTA ACA TGG GGC GTC TTA GCT TTA TCA GGC CGA TTA TCG TCG TCG TCC
 ASN OPA Leu Asp Val ASN Thr Leu ASN Lys Ala Trp Thr Tyr Leu Thr Ser Gly Val Leu Ala Leu Leu Gly Arg Leu Ser Ser Ser Ser
 6121/2041 6151/2051 6181/2061
 CAA CCT TCG TCG TTA GAA GTT GCT TCC GAA GAC GAT TTT GCC ATA GCC ACR CGA CGC CTC TTA ACT GTC TCG CCT AAC ACG TCC GCG ATC
 Glu Pro Ser Ser Leu Glu Val Ala Ser Glu Esp Asp Phe Ala Ile Ala Thr Arg Arg Leu Leu Ile Val Ser Ala ASN Thr Ser Ala Ile
 6211/2071 6241/2081 6271/2091
 AAA ATT GTC GTT GAG CTT TTT GGA ATT ATT TCT GAT TGC CGG CCT TTT TGG GCG GGT TTC ATT CTA ACT GTG CCC GAT TTT ATT TCA GAC
 Lys Phe Val Val Glu Leu Phe Gly Ile Ile Ser Asp Cys Gly Arg Phe Trp Ala Gly Phe ASN Leu Thr Val Pro Asp Phe ASN Ser Asp
 6301/2101 6331/2111 6361/2121
 AAC ACG TTA GAA AGC GAT GGT GCA GGC CCT GGT AAC ATT TCA GAC GCC AAA TCT ACT ATT GGC CCT CCT GGT GAT GAT AAA TCT
 Asp Thr Leu Glu Ser Asp Gly Ala Gly Gly Cys ASN Ile Ser Asp Gly Lys Ser Thr ASN Gly Gly Gly Gly Ala Asp Asp Lys Ser
 6391/2131 6421/2141 6451/2151
 ACC ATC GGT GGA CGC CCA CGC GGG CCT GGC GGC GCA CGC ASN CCT CCT GGC CCT GAT GCA GAC GGC GGT TCA GGC TCA ATT GTC
 Thr Ile Gly Gly Gly Ala Gly Gly ASN Gly Gly Gly Gly Gly Gly Gly Gly Ser ASN Asp Gly Gly Leu Gly Ser ASN Val
 6481/2161 6511/2171 6541/2181
 TCT TTA GGC AAC ACA GTC GGC ACC TCA ACT ATT GTC CTG GTC TCC GGC CCT GGT TTT GGT ATT ACC GGT CTG AGA CGA CTG CGA TTT
 Ser ASN Ile Gly ASN Thr Val Gly Thr Ser Thr Ile Val Leu Val Ser Gly Ala Val Phe Gly Leu Thr Gly Leu Arg Val Arg Phe Phe
 6571/2191 6601/2201 6631/2211
 TCG TTT CTA ATA GCT TCC AAC ATT TGT TGT CTG TCG TCT AAA GGT GCA GCG CCT TCA CGT TCC GTC CGC ATT CCT GGA CGC CGC ATT
 Ser Phe Leu Ile Ala Ser ASN ASN Cys Cys Leu Ser Ser Lys Gly Ala Ala Gly OPA Gly Ser Val Gly Ile Gly Gly Ala Gly ASN
 6691/2221 6691/2231 6721/2241
 TCA GAC ATC GAT GGT GGT GGT GCA GGC CCT GGA ATG TAA GGC ACC GGA GAA CCT GGT GGC CGC CGC CGC CGT ATA ATT TGT
 Ser Asp Ile Asp Gly Gly Gly Gly Ala Gly Met Leu Gly Thr Gly Glu Gly Gly Gly Gly Ala Ile Ile Cys 6751/2251 6781/2261 6811/2271
 TCT GGT TTA GTT TGT TCG CGC ACG ATT STG GGC ACC GGC GCA GGC CCT GGT GGC TGC ACA ATT GCA GGT CCT CGT CTC ATT CGA CGC AGC CCT
 Ser Gly Leu Val Cys Ser Arg Thr Ile Val Gly Thr Gly Ala Gly Ala Gly Cys Thr Thr Glu Gly Arg Leu Arg Gly Ser Ala
 6841/2281 6871/2291 6901/2301
 TGG GGT GGT GGC AAT TCA ATA TTA TAA TGG GAA TAC AAA TCG TAA AAA TCT GGT ATA AGC ATT CTA ATT TCG CTA TCG TTT ATT ACC GTG CGG
 Trp Gly Gly ASN Ser Ile Leu OCH Leu Glu Tyr Lys Ser OCH Lys Ser Ala Ile Ser Ile Val Ile Ser Leu Ser Phe Thr Val Pro
 6931/2311 6961/2321 6991/2331
 ATA ATT AAC AAC CGC TCA ATT TAA GCA ATT GTC TIG TAA ATA GAT TGT CTC CCT CCT GGT CGG ATA ACA AGC CTT TTC ATT TTT ACT
 Ile Phe ASN ASN Arg Ser Met OCH Ala Ile Val Leu OCH Arg Asp Asp Cys Leu Lys Leu Arg Thr Pro Ile Thr Ser Leu Phe Ile Phe Thr
 7021/2341 7051/2351 7081/2361
 ACA GCA TTG TAG TGG CGA GAC ACT TCG CTG TCG TCG ACC TAC ATT TGT CCT TGG TTG TCA AAA AGC TCG TTG CGC AGC CCT AAA ATA ATT
 Thr Ala Leu AMB Trp Arg Asp Thr Ser Leu Ser Thr Val Leu Ser OCH Met Leu Phe Leu Ile Ile Cys ASN ASN Ile Val Ser Thr Arg Ser Lys ASN
 7111/2371 7141/2381 7171/2391
 AAA AGA ACA TCT CGT TPC ACC ACC ACT GTC TTG TCG TAA AGG TTG ATT TGG ATT TGG CCT TCC GCA GTC TCG AAC CGT TCA AAA ATT
 Lys Arg Thr Ser Leu Phe Ser Thr Val Leu Ser OCH Met Leu Phe Leu Ile Ile Cys ASN ASN Ile Val Ser Thr Arg Ser Lys ASN
 7201/2401 7221/2411 7261/2421
 TGA TCC GCA TCA ATT TTG TTG TTC CTA TTA TGG ATT AAA TAA GAT TGT ACA GAT TCA ATT CTA CGA TTC GTC ATT GGC ACC ACC AAC ATT CCT
 OPA Cys ASN Ile Ser Ile Leu Leu Phe Leu Ile ASN Cys ASN Asp Cys Thr Asp Ser Tyr Leu Arg Phe Val Met Ala Thr Thr ASN Ala
 7291/2431 7321/2441 7351/2451
 ACG CCT CAA AGC CGT CTA CAA ATT TAC GAA AAC TGC AAA AAC ACT CGG TAT AAA ATA ATT AAC GAC GGC CGC TTT GGC AAA ATA ATT CCT
 Thr Leu Glu ASN Thr Leu Val Glu ASN Phe Tyr Glu ASN Cys Lys ASN Val Lys Thr Arg Tyr Lys Ile Ile ASN Gly Arg Phe Gly Lys Ile Ser
 7381/2451 7411/2471 7441/2481
 ATT TTA TCG CAC AAG CCC ACT ACC AAA ATT TGT ATT STG CAG AAA ACA ATT TCG GCG CAC ATT ATT AAC GCT GAC GAA ATA AAA ATT CAC CAG
 Ile Leu Ser His Lys Pro Thr Ser Lys Leu Tyr Leu Glu Lys Thr Ile Ser Ala His ASN Phe ASN Ala Asp Glu Ile Lys Val His Glu
 7471/2491 7501/2501 7531/2511
 TTA ATG AGC GAC CAC CCA ATT ATT ATA AAA ATT TAT ATT AAC CAC GGT TCC ATT AAC GAC CAA ATT GTC ATT GTC ATT GTC ATT GAC TAC ATT GAC TGT
 Leu Met Ser Asp His Pro ASN Phe Ile Lys Ile Tyr Phe ASN His Gly Ser Ile ASN ASN Glu Val Ile Val Met Asp Tyr Ile Asp Cys
 7561/2521 7591/2531 7621/2541
 CGC GAT TTA ATT GAA ACA CTA CAA ATT AAA GGC GAG CTT TCG TAC CAA ATT GTC ATT AAC GAG CAG CTG TGT GAA GGC CTC AAC
 Pro Asp Leu Phe Glu Thr Leu Glu Ile Lys Gly Glu Leu Ser Tyr Glu Leu Val Set ASN Ile Arg ASN Glu Ile Leu Cys Glu Ala Leu ASN

pACgp67-ScFv461 -> 1-phase Translation

Fig. 1D

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7651/2551 7661/2561 7711/2571
 GAT TTG CAC AAG CAC AAT TTC AEA CAC AAC GAC ATA CAA CTC GAA AAT GTC TTA TAT TTC GAA GCA CTT GAT CGC GTG TAT GTT TGC GAT
 asp leu his lys his asn phe ile his asn asp ile lys leu glu asn val leu tyr phe glu ala leu asp arg val tyr val cys asp
 7741/2581 7771/2591 7801/2501
 TAC GGA TTG TGC AAA CAC GAA AAC TCA CTT AGC GTG CAC GAC CGC ACG TTG GAG DAT TTT AGT CGG GAA AAA ATT CGA CAC ACA ACT ACT
 tyr sly leu cys lys his glu asn ser leu ser val his asp gly thr leu glu tyr phe ser pro glu lys ile arg his thr thr met
 7831/2611 7861/2621 7891/2531
 CAC GTT TCG TTT GAC TGG TAC CGG CGG TGT TAA CAC ACA ACT TCC TAA CGG GCG gct cgt ATT CAT GGT CAT AGC TGT TTC CTG TGT GAA
 his val ser phe asp trp tyr ala ala cys OCH his thr ser cys OCH pro ala val arg asn his gly his ser cys phe leu cys glu
 7921/2641 7951/2551 7981/2651
 ATT GTT ATC CGC TCA CAA TTC CAC ACA ACA TAC GAG CGG GAA GCA TAA AGT GTC AAC CCT GGG GTG CCT ATT GAG TGA GCT AAC TCA CAT
 ile val ile arg ser glu phe his thr thr tyr glu pro glu ala OCH ser val lys pro gly val pro asn glu OPA ala asn ser his
 8011/2671 8041/2581 8071/2691
 TAA TTG CGT TGC GCT CAC TCC CGG CTT AGT CGG GAA ACC TGT CGT GGC AGC TGC ATT ATT GAA TCG GCC AAC GCG CGG GGA GAG GCG
 OCH leu arg cys ala his cys pro leu ser ser arg glu thr cys arg ala ser cys ile asn glu ser ala asn ala arg gly glu ala
 8101/2701 8131/2711 8161/2721
 GTT CGT TCA TTG GGC GCT CTT CGG CTT CCT CGC TCA CTG ACT CGC TGC GCT CGG TCG TGT GGC TGC CGC CAG CGG TAT CAG CTC ACT CAA
 val cys val leu sly ala leu pro leu pro arg ser leu thr arg cys ala arg ser phe gly cys gly glu arg tyr glu leu thr glu
 8191/2731 8221/2741 8251/2751
 AGG CGG TAA TAC GGT TAT CCA CAG PAA AGC CAG CAG GAA ASA ACA TGT GAG CAA AAG GCC AGC AAA AGG CCA GGA ACC GTC AAA
 arg arg OCH tyr gly tyr pro glu asn glu ile thr glu glu arg thr cys glu glu lys ala ser lys arg pro gly thr val lys
 8281/2761 8311/2771 8341/2781
 AGG CGG CGT TGC TGG CGT TTT TCC ATA GGC TCC GCC CCC CTG ACG AGC ATC ACA AAA ATT GAC GCT CAA GTC ACA GGT CGC GAA ACC CGA
 arg pro arg cys trp arg phe ser ile gly ala pro leu thr ser ile thr lys ile asp ala glu val arg gly glu thr arg
 8371/2791 8401/2801 8431/2811
 CAG GAC TAT AAA GAT ACC AGG CGT TTC CCT CTG GAA GCT CCC TCG TCC GCT CTC CTG TCC CGA CCC CGC TTA CGG GAT ACC TGT CGG
 glu asp tyr lys asp thr arg arg phe pro leu glu ala pro ser cys ala leu leu phe arg pro cys arg leu pro asp thr cys pro
 8461/2821 8491/2831 8521/2841
 CCT TTC TCC CTT CGG GAA CGG TGG CGC TTT CTC ATA GCT CAC GCT GTC GGT ATT TCA GTT CGG TGT AGG TCG TTC GCT CCA ACC TGG CCT
 pro phe ser leu arg glu ala trp arg phe leu ile ala his ala val gly ile ser val arg cys arg ser phe ala pro ser trp ala
 8551/2851 8581/2861 8611/2871
 GTG TGC ACG AAC CCC CGG TTC AGC CGG ACT GCT CGG CCT TAT CGG GTC ACT ATT CGT CCT TGT AGT CCA ACC CGG TAA GAC ACG ACT ATT CCT
 val cys thr asp pro pro phe ser pro thr ala ala pro tyr pro val thr ile val leu ser pro thr arg OCH asp thr thr tyr arg
 8641/2881 8671/2891 8701/2901
 AAC TGG CAG CAG CCA CTG GDA ACA GGA TTA CGA CGG GGT ATT TAG CGG GTG CTA CGA ATT TCT TGA ACT GGT CGC CTA ACT ACG CCT
 his trp glu glu pro leu val thr gly leu ala glu arg gly met AMB ala val leu glu ser ser OPA ser gly gly leu thr ala
 8731/2911 8761/2921 8791/2931
 ACA CTA GAA GGA CAG TAT TTG GTC TCT CGG CTC TGC AGC CAG TAA CCT TCC GAA AAA GAG TTG GTC GCT CCT GAT CGG GCA AAC AAA
 thr leu glu gly leu ser ala leu cys OPA ser glu leu pro ser glu lys glu leu val ala leu asp pro ala asn lys
 8821/2941 8851/2951 8881/2961
 CGA CGG CTG GIA CGG GTG TTT TTG GTC AAC ACC AGA TTA CGC GCA GAA AAA AAG GAT CTC AAC AAG ATT CCT TGA TCT TTT CTA
 pro phe leu val ala val val phe leu phe ala ser ser arg leu arg ala glu lys lys asp leu lys lys ile leu OPA ser phe leu
 8911/2971 8941/2981 8971/2991
 CGG CGT CTG ACG CTC AGT CGG ACT AAA ATT AAC CTC TTG AAG GGA TTT TTG TCA TGA GAT TAT CAA AAA GGA TCT TCA CCT AGA TCC TTT TAA
 arg gly leu thr leu ser gly thr lys thr his val lys gly phe trp ser OPA asp tyr cys glu lys gly ser ser pro arg ser phe OCH
 9001/3001 9031/3011 9061/3021
 ATT AAA AAT GAA GTT TTA ATT CAA TCT AAA GTC TAT ATG AGT AAA CCTT GGT CTG ACA ATT ATT ATT CCTT GCT TAA TCA GTC AGG CAC CTA CCT
 ile lys asp glu val leu asp glu ser lys val tyr met ser lys leu gly leu thr val the asp ala OCH ser val arg his leu ser
 9091/3031 9121/3041 9151/3051
 CAG CGA CCT GTC TAT TTG GTC CAT CGA TAG TTG CGT GAC TCC CGG TCG TGT AGA TAA CTA CGA TAC CGG AGG CCT TAC CAT CTG CGC CGA
 glu arg ser val tyr phe val his pro AMB leu pro esp ser pro ser cys arg OCH leu arg tyr gly arg ala tyr his leu ala pro
 9181/3051 9211/3071 9241/3081
 GTG TTG CAA TGA TAC CGC GAG ACC CAC CCT CAC CGG CTC CAG ATT TAT CAG CAA TAA ACC AGC CAG CGG CGG CGC AGC GCA GAA GTG
 val leu glu OPA tyr arg glu thr his ala his arg leu glu ile tyr glu OCH thr ser glu pro glu gly pro ser ala glu val
 9271/3091 9301/3101 9331/3111
 GTC CGT CAA CCT TAT CGG CCT CGA TCC ACT CTA TTA ATT CCT GCG AAG CTC GAC TAA GTC GGT CGC CAG TTA ATA CCTT TGC CGA ACC
 val leu glu leu tyr pro pro phe ser ser leu leu ile val ala gly lys leu glu OCH val val arg glu leu ile val cys ala thr
 9361/3121 9391/3131 9421/3141
 TTG TTG CCA TTG CTA CAG GCA TCG TGG TGT CAC GCT CGT COT TTG GTC TGG CCTT CAT TCA GCT CGT CGG CCT CCC AAC GAT CAA CGC GAG TTA
 leu leu pro leu leu glu ala ser trp cys his ala ala arg arg leu val trp leu his ser ala pro val pro asp asp glu gly leu
 9451/3151 9481/3161 9511/3171
 CAT GAT CCC CCA TGT TGT GCA AAA AAG CGG TTA CCT CCT CGC GTC CTC CGA TGG TTG TCA GAA GTC AGT TGG CGG CAG TGT TAT CAC TCA
 his asp pro pro cys cys ala lys lys arg leu ala pro ser val leu arg ser leu ser glu val ser trp pro glu cys tyr his ser
 9541/3181 9571/3191 9601/3201
 TCG TTA TGG CAG CAC TCC ATA ATT CTC TTA CTG TCA TGC CAT CGG CAA GAC GCT TTT CGT AGT ACT CAA CCA ACT CAT CCT
 trp leu trp glu his cys ile ile leu leu ser cys his pro OCH esp ala phe leu OPA leu val ser the glu pro ser his ser
 9631/3211 9661/3221 9691/3231
 GAG AAT AGT GTC CGC GAC CGA CCT TGT CCT CGG CGT CAA TAC CGG ATA ATA CGC CAC ATA GCA GAA CCT TAA AAG TGC CGA
 glu asp ser val cys gly esp arg val ala leu ala arg arg glu lys gly ile ile pro arg his ile ala glu leu OCH lys cys ser
 9721/3241 9751/3251 9781/3261
 TCA TTG GAA AAC GTT CCT CGG CGC GAA AAC TCT CAA GGA TCT TAC CGC TGT TAA GAT CGA CCT TTG CGT TGT AAC CCA CTC GTG CAC CGA ACT
 ser leu glu asp val leu arg gly glu asp ser glu gly ser tyr arg cys OPA esp pro val arg cys asp pro leu val his pro thr
 9811/3271 9841/3281 9871/3291
 GAT CCT CAG CAT CCT TTA CCT TCA CCA CGG TTT CTG GGT GAG CAA AAA CAG GAA CGC AAA ATT CGC CAA AAA AGG GAA TAA GGG CGA CAC
 esp leu glu his leu leu leu ser pro ala phe leu gly glu glu lys glu glu gly lys met pro glu lys arg glu OCH gly arg his
 9901/3301 9931/3311 9961/3321
 GGA AAC GTT GAA TAC TCA TAC TGT TCC TTT TCC ATT ATT ATT GAA CGA TTT ATT AGC GGT ATT GTC TCA TGA CGG GAT ACA TAT TTG ATT
 gly asp val glu tyr ser tyr ser ser phe phe asp ile ile glu ala phe ile arg val ile val ser OPA ala esp thr tyr leu asp
 9991/3331 10021/3341 10051/3351
 GTC CCT ATA AAA ATA AAC AAA TAG GGG TTC CGC CGA CAT TCC CCC GAA AGG TGC CAC CTG ACG CCT ATT AAG AAA CCA TAA TCA TGA CAT
 val phe arg lys ile asp lys AMB gly phe arg ala his phe pro glu lys cys his leu the ser lys pro leu leu ser OPA his
 10081/3361 10111/3371 10141/3381
 CAA CCT ATA AAA ATA CGC CGT TCA CGA CGC CCT TTC GTC TCG CGC GFT TCG CCT ATT AGC CGC AAA ACC CCT GAC ACA TGC ACC TCC CGG
 CGC pro ile lys ile gly val ser arg gly pro the val ser arg val ser val met thr val lys thr ser asp thr cys ser ser arg
 10171/3391 10201/3401 10231/3411
 AGA CGG TCA CAG CCT GTC TGT AAG CGG ACT CGG GGA CGA GAC AAG CGC CGT AGG CGG CGT CGG CGG GTG TTG CGG GGT GTC CGG CCT CGC

pACgp67-ScFv461 -> 1-phase Translation

Fig 1E

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10261/3421 10291/3421 10321/3441
 TTA ACT ATG CGG CAT CAG AGC AGA TTG TAC TGA GAG TGC ACT ATA TGC GGT GTG AAA TAC CGC AGA GAT CGG TAA GGA GAA AAC ACC GCA
 leu thr met arg his gln ser arg leu tyr CTA glu cys thr ile cys gly val lys tyr arg thr asp ala CCA gly glu arg thr ala
 10351/3451 10381/3461 10411/3471
 TCA GGC GCC ATT CGC CAT TCA GGC TGC CGA ACT GTT GGG AAG GGC GAT CGG TGC CGG CCT CTT CGG TAT TAC CGC AGG TGG CGA AAC CGG
 ser gly ala ile arg his ser tyr cys ala thr val gly lys gly asp arg cys gly pro leu arg tyr tyr ala ser trp arg lys gly
 10441/3481 10471/3491 10501/3501
 GAT GTG CTG CAA CGC GAT TAA GTT GGG TAA CGC CAG GGT TTT CCC AGT CGC GAC GTT GTA AAA CGA CGG CCA GTG CC
 asp val leu gln gly asp CCA val gly phe pro ser his asp val val lys arg arg pro val

*** DNA Strider™ 1.2 *** Mardi 16 janvier 2001 21:58:26

pACgp67-ScFv350 -> 1-phase Translation

DNA sequence 10511 b.p. AAGCTTATACCG ... AGGGCTATGCG line 1

1/1 31/11 61/21
AAG CTT TAC TCG TAA AGC GAG TTG AAG GAT CAT ATT TAG TCG CGT TTA TGA GAT AAG ATT GAA AGC ACG TGT AAA ATG TTT CCC GCG CGT
lys leu tyt ser OCH ser glu leu lys asp his ile AMB leu arg leu OPA arg lys ile glu ser thr cys lys met phe pro ala arg
91/31 12/1/41 151/51
TGG CAC AAC TAT TTA CAA TCG GGC CAA GTT ATA AAA GAT TCT AAT CTG ATA TGT TTT AAA ACA CCT TIG CGG CCC GAG TIG TTT CGG TAC
trp his ASN tyt leu glu cys gly glu val ile lys asp ser ASN leu ile cys phe lys the pro leu arg pro glu leu phe ala tyt
18/1/61 21/1/71 24/1/81
GTG ACT ACC GAA GAA GAT GTG TGG ACC GCA GAA CAG ATA GIA AAA CAA AAC CCT ACT ATT GGA GCA ATA ATC GAT TTA ACC AAC ACG TCT
val thr ser glu glu asp val trp thr ala glu glu ile val lys glu ASN pro ser ile gly ala ile ile asp leu thr ASN thr ser
27/1/91 30/1/101 33/1/111
AAA TAT TAT GAT GGT GTG CAT TTT TTG CGG CCC GGC CTG TTA TAC AAA AAA ATT CAA GCA CCT GGC CGG ACT TTG CGG CCT GAA AGC ACA
lys tyt tyt asp sly val his phe leu arg ala gly leu leu tyt lys ile glu val pro gly glu thr leu pro pro glu ser ile
35/1/121 39/1/131 42/1/141
GTT CAA GAA TTT ATT GAC ACG GEA AAA GAA TTT ACA GAA AAC TGT CCT CGC ATG TTG CGT GGC GTG CAC TGC ACA CAC GGT ATT AAT CGC
val glu glu phe ile asp thr val lys glu phe thr glu lys cys pro gly met leu val gly val his cys thr his gly ile ASN arg
45/1/151 48/1/161 51/1/171
ACC GGT TAC ATG GTG TGC AGA TAT TTA ATG CAC ACT CTG GGT ATT CGG CGG CAG GAA GGC ATA GAT AGA TTC GAA AAA GGT AGA GGT CAC
thr gly tyt met val cys arg tyt leu met his thr leu sly ile ala pro glu ala ile asp arg phe glu lys ala arg gly his
54/1/181 57/1/191 50/1/201
AAA ATT GAA AGA CAA AAT TAC GTT CAA GAT TTA TIA ATT TAA TTA ATA TTA TTT GCA TTC TTT AAC AAA TAC TTT ATC CTA TTT TCA AAT
lys ile glu arg glu ASN tyt val glu asp leu leu ile leu phe ala phe the asp arg phe glu lys tyt phe ile leu phe ser ASN
63/1/211 65/1/221 68/1/231
TGT TGC GCT TCT TCC AGC GAA CCA AAA CTA TGC TTC GGT TGC TCT TGT CCT TAC GGT GCA GAT CAG TGC CCT TGT CGG TAC
cys cys ala ser ser ser glu pro lys leu cys phe ala cys ser val AMB leu val ala 78/1/261
72/1/241 75/1/251
GAT TAG GCC TAT TCT CCA CCA CAA TGT TGG CAA CGT TGA TGT TAC GGT ATT CCT TAT GGT TTT GGT TTA CCA CCT ACT CGC TCT TTT CGC CGG TAA
asp AMB ala gly tyt ser pro pro glu cys trp glu arg OPA cys tyt val tyt ala phe 87/1/291
81/1/271 84/1/281
TAG CGG TAA AGC TAG TCC COT CGC CGG TCA CGC ACA ACA CGG GAT GGT TGC GGT CCT CGG CGG GGT ATT GAA CGG CGC GAT CGG ACA AAT
AMB pro OCH thr AMB cys arg arg ala ser arg thr pro asp val cys ala cys pro arg gly ile glu pro arg asp pro thr ASN
90/1/301 93/1/311 96/1/321
CCA CCA CCT TCG CAA CTA AAT CGG TGA CCT GGG CGT CCT TTT TGT CCT TAA TTT CCT CTT TGT TCA TGG TTT CCT CGG TAC
pro pro leu trp glu leu ASN arg OPA pro ala arg leu phe ser ala leu phe arg leu ser the ala trp phe pro gly ser arg cys
99/1/331 102/1/341 105/1/351
ACA TGC GGT TTA GAT CAG TCA TGA CGC CGG TGA CCT GCA AAC CCT TGG CCT CGA TGT CCT GGT ATT GAA CGG CGC GAT CGG ACA AAT
thr cys gly leu asp glu ser OPA arg ala OPA pro ala ASN leu trp pro arg ser ala 114/1/381
108/1/361 111/1/371
ACT CCT GTT TTT TAA CAA GTT CCT CGG TTT TGT CCT CCA CGG CCT TGT GCA CGG CCT TGT CCT GGT CCT CGG TAC TCA CCT TAG
the leu val phe OCH glu val pro arg phe ala pro pro pro leu ala ala arg leu cys ala arg CPA met ser glu ser ala AMB
117/1/391 120/1/401 123/1/411
TCA CCA ACT GTT TGC TCT CCT CCT CCT GGT TGA TGG CGG GAT CCT ACT CCT CGG TGC GGT ATT GAA CGG CCA CCT GAA TIA CCT CCT CTA AAA
ser pro thr val cys ser pro pro pro val val OPA ser arg arg thr cys arg cys 126/1/421 129/1/431 132/1/441
CGC CCT CCT GTA ATT CTA CGG CCT AAC GCA ATT TGG ACT TCA CAA TCA CGT GAA TCA CGG CGC ATT TAC TAA TGA CCA CCT TGT CCT GGT
ala ile leu val ile leu trp arg lys ala ile trp thr ser OCH ser ala glu ser arg 135/1/451 138/1/461 141/1/471
CCA ATT ACA CGG GGT CGC CCC TTT TCA CGA CGC TGT TAC AGG TAG CGC CGG CAT TTT GCA TGG CCT CAA ATA ACC ATT TGT ATT TAT
ala ASN thr ala gly arg pro phe ser arg arg cys AMB arg AMB gly pro his phe sly 144/1/481 147/1/491 150/1/501
TGT CCA CAT GAA CAC CTA TAC CCT IAT CAC AAA CCT IAT ATT TIA AAC TGT TAG CGA CGT CCT TGG CCA CGG ACT GGA CCT GTT CCT CGC
cys leu his glu his val AMB leu tyt his lys leu tyt ile leu ASN cys AMB arg arg pro pro pro arg thr gly pro val sly arg
153/1/511 156/1/521 159/1/531
GCT CTA CGA CGT ACC GCA CGT TGA AGC TAT CCT CTC CAA ATT TAA ATT CTC CAA TTT TAA CGC GAG CCA TTT TGA TAC ACC TGT GTC GAT
ala leu ala arg thr ala gly OPA thr tyt leu leu glu ile OCH ile leu glu phe OCH 162/1/541 165/1/551 168/1/561
CTT TCA ACA ACT ATT GTT TTT TAA CGC AAA CTA AAC TTA TTG TGG TAA GCA ATA ATT AAA ATT CGG GGA ACA TGC CGC CCT ACA ACA CTC
phe ala thr thr ile val phe OCH arg lys leu ASN leu leu trp CGH ala ile ile lys 171/1/571 174/1/581 177/1/591
CTC GTC ATT AAC GCA GAC CGC CGG CGT CCT CGC GCA AGC CGC CAA AAC GTG TTG CGC CCT CAA CGC CGC AAA CAT CGC AAA ACC CAA TAG
val val met ASN ala asp gly ala gly leu gly ala ser gly OCH ASN val leu arg val glu arg gly lys his arg lys ser glu AMB
180/1/581 183/1/591 185/1/621
TAC ACT ATT GAT TTG CAT ATT AAC CGC GAT TTT TTA ATT TAT CCT ATT TAA TAA ATA ATT ATG ACG CCT ACA ACT CCC CGC CGG CGT TGA
tyt ser phe asp leu his ile ASN gly asp phe leu ASN tyt leu ile OCH CGH ile val met thr pro thr thr pro arg pro arg OPA
188/1/631 192/1/641 195/1/651
CTC CCT CGA CCT CGA CGA GGT CCT CGT TGA CGC CCT CCT CGC TGT CGC CAA ACA CGT CGA CGC CGT CGT CGA TGA CCA CGG CGG TCC CGC ACT
leu ala ala pro arg ala val arg OPA arg leu pro pro cys gly arg thr arg arg ala 198/1/651 202/1/671 204/1/681
CGA CGC ACA AGT ATC TGT ACA CGC AAA GAT CCT CGG CGC AAG GCA CGT CGG CGT CCT CGA AGT GGC AAT ATT CGC AAA TTC CAA AAT ATA TAC
arg arg thr ser ile cys thr pro ASN asp arg arg ala lys ala arg arg pro pro ser gly ASN ile gly lys phe glu ASN ile tyt
207/1/691 210/1/701 213/1/711
AGT TGG CCT GTT TGC GCA ATT CTA TCG TGG CGT TGG GCA TGT ACC TCC GAA CGT TGA TTT CGA TGC AAG CGG AAA TIA AAT CAT TGC GAT
ser trp val val cys ala tyt leu ser trp arg trp ala cys thr ser glu arg OPA phe ala cys lys pro lys leu ASN his cys asp
216/1/721 219/1/731 222/1/741
TAG TGC GAT TAA AAC GTT CTA CAT CCT CGC TTT TAA TCA TGC CGT CGA TTA ATT CGC CGA ATG GAG TCA ATG GAT CAA AGT GTG GAA TAA
AMB cys asp OCH ASN val val his pro arg phe OCH ser cys arg arg leu ASN arg ala ile glu ser ser ASN glu ser val glu OCH
225/1/751 228/1/761 231/1/771
TGT CCT GTT TGT ACC CCC GAG TCA AGC GCA CGC CGT ATT TTA ACA AAC TAG CGA CCT TGT TGT AGG TIA GTT TTT AAT GCA ACT TTA CGC
lys the leu cys ile pro glu ser ser ala arg ile leu ASN AMB pro ser cys lys leu val ser phe ASN ala thr leu ser
234/1/781 237/1/791 240/1/801

Fig 2

pACgp67-ScFv350 -> 1-phase Translation

Fig. 2B

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2431/811 2461/821 2491/831
 AAT TAA ATA CCT TGC GAC GCA ACG TGC ACG ATC TGT GCA CGC GTC CCG GCA CGA GCT TGC ATT GCA ATA ACT TTT TAC GAA GCG ATG ACG
 asn OCH ile ala cys asp ala thr cys thr ile cys ala arg val pro ala arg ala leu ile val ile ser phe tyt glu ala met thr
 2521/841 2551/851 2581/861
 TGA CCC CGG TAC TGA CAA CGA TCA CGC CCA AAA GAA CTG CGT ACT ACA AAA TTA CGG AGT ATG TCG GTG AGC TTA AAA CTA TCA AGC CTC
 OPA pro pro AMB OPA gln arg ser arg pro lys glu leu pro thr thr lys leu pro ser ser val thr leu lys leu leu ser his
 2611/871 2641/881 2671/891
 CCA ATC GAC CGT TAG TGC AAT CAG GAC CGC TGC TGC GAG AAG CGG CGA AGT ATG CGG AAT GCA TCG TAT AAC GTG TGG AGC CCG CTC ATT
 pro ile asp arg AMB ser asn gln asp arg 2701/901 2731/911 2761/921
 AGA GCG TCA TGT TTA GAC AAG AAA GCT ACA ATT TCA ATT GAT CCC GAT GAT TTT ACT GAT AAA TTC ACC CTA ACT CCA TAC ACC GTC TTC
 arg ala ser cys leu asp lys ala thr tyt leu ile asp pro asp phe ile asp lys leu thr leu thr pro tyt thr val phe
 2791/931 2821/941 2851/951
 TAC AAT GGC GGG GTT TTG GTC AAA ATT TCC GCA CTG CGA TTG TAC ATG CTG TTA ACG GCT CGG CCC ACT ATT AAT GAA ATT AAA ATT TCC
 tyt asn gly gly val leu val lys ile ser gly leu arg leu tyt met leu leu thr ala pro pro thr ile asn glu ile lys asn ser
 2881/961 2911/971 2941/981
 AAT ATT AAA AAA CGC AGC AAG AGA AAC ATT TGT ATG AAA GAA TGC GTC GAA GCA AAG AAA ATT GTC GTC AAC ATG CGC AAC AAC AAG ATT
 asn phe lys lys arg ser lys arg asn ile 3001/1001 3031/1011
 AAT ATG CCT CCG TGT ATA AAA ATA TTG AAC GAT TTG AAA GAA AAC ATT GTC CGG CGC GGT ATG TAC AGG AAC AGG TTT ATA CTA
 asn met pro pro cys ile lys ile leu 3061/1021 3091/1031 3121/1041
 AAC TGT TAC ATT GCA AAC GTG GTT TCG TGT GCC AAC TGT GAA AAC CGA TGT TTA ATC AAC GCT CTC ACC CAT TTC TAC AAC CAC GAC TCC
 asn cys tyt ile ala asn val val ser cys 3151/1051 3181/1061 3211/1071
 AGG TGT GTG CGT GAA GTC ATG CAT CTT TTA ATC AAA TCC CAA GAT GTG TAT AAA CCA CCA AAC TGC CAA AAA ATG GAC AAC lys
 lys cys val gly glu val met his leu leu 3241/1081 3271/1091 3301/1101
 CTC TGT CGG TTT GCT GGC AAC TGC AAG GGT CTC ATT CCT ATT TGT ATT TAT TGA ATA ATA AAA CAA TTA TAA ATG CTC ATT TIG TTT TTT
 leu cys pro phe ala sly asn cys lys gly 3331/1111 3361/1121 3391/1131
 ATT AAC GAT ACA AAC CAA ACG CAA CAA GAA CAT TTG TAG TAT TAT CTC TAA TTA TTS AAA ACC CGT ACT ATT AAT CCC TGA GGT ATT ATT TAA
 ile asn asp thr asn gln thr gln gln 3421/1141 3451/1151 3481/1161
 AAT CAT ATT CAA ATG ATT CAC AGT TAA TTT CGG ACA ATA TAA TTT TAT TTT CAC ATA AAC TAG AAC CCT TGT CGT CTT CTT CGT ATT
 asn his phe gln met ile his ser OCH phe 3511/1171 3541/1181 3571/1191
 CCT TGT CTT TTT CAT ATT CCT CAT AAA ATT TAA CAT ACT ATT TAT CGT ATC CAT ATA TGT ATT ATT CGT ATA GAG TAA ATT TTT TGT
 pro ser leu phe his ser pro his lys 3561/1201 3631/1211 3661/1221
 TGT CAT AAA ATT ATA TGT CTT TTT TAA TCG ATT GTA TAG TAC CGC TGC GCA TGC TTT TGC ATT ATT TAA CAA CAG CGC TAT ATT CTC GTC
 cys his lys tyt ile cys leu phe OCH trp 3691/1231 3721/1241 3751/1251
 GTT CTT CGG ACT GTG TTG CTT TAA CTA CTA ATT TAA TAT CAA TGA ATT TGG GAT CGT CGG TTT TGT ACA ACA TGT TGC CGG CAT ACT
 val leu arg ser val leu leu OCH leu leu 3781/1261 3811/1271 3841/1281
 ACT CAG CTT CTT CTC CTT CAA TTA CGC CAT ATT TGA CGA GCA CGG GAT TAA CGC AAC ATT CCT CCA AAA TGT TTT ACT AAC CGT TAA ACA AAA
 thr gln leu leu val gln leu his his phe leu ala ala pro esp OCH his asn phe 3871/1291 3901/1301 3931/1311
 ACA GTT CAC CTC CCT TTT CTC TAC ATT TGT CGT CGA CGA ATT TGT TGT TAA AAA TAA TAA CAG CCA TTG TAA TGA GAC GCA ACT ATT
 thr val his leu pro phe leu tyt tyt cys leu arg ala val val cys cys OCH lys OCH 3961/1321 3991/1331 4021/1341
 ATC ACA AAC TGG AAA TGT CTC TCA ATA ATT ACT TGC TGA TAC CAT CGA GAT ATT TAA ATT ACT AAC CAT CTC CGA ATT AAA TAA GTC ATT
 ile thr asn trp lys cys leu ser ile tyt ser cys CGA tyt his gln asp asn OCH asn 4051/1351 4081/1361 4111/1371
 TAC CGT TTT CGT AAC AGT ATT GTC ATA AAA AAA CCT ATA ATT ATT CGG GAT ATT TCA TAC CGT CCC ACC ATT CGG CGC CGA TGT CTC ATT
 tyt cys phe arg asn ser phe val ile lys 4141/1381 4171/1391 4201/1401
 CTC GCA ATT CAG TCA CAC CAA CGC TTC ATT AAG GAA CAC ACA ACC AAC AAG ATG GTC AGC GCT ATT ATT ATT ATT ATT ATT ATT ATT ATT
 leu val asn gln ser his gln gln phe asn his thr ser lys met val ser ala ile val leu tyt val leu leu ala ala ala 4231/1411
 4261/1421 Tag HIS Factor Xa 4291/1431 → ScFv 350 = VH 4381/1461
 CGG CGT TGT GCG TTT CGC CGG GAT CTT gga tcc CAT CAT CAC CAC CAC CGC att gaa gga aga GCA TTG CAG GTC CAA CTG CAG CAG TCT
 ala his ser ala phe ala Val asp leu gln ser his his his his ile glu gln 4331/1441 4351/1451 4381/1461
 GGG GCT GAA CTG GCA AAA CCT CGG CGC TCA GTC AAG CGT TGT CGC CAC ACC ATT ACT ACC TAC TGG ATT CGC CGC TGG GCA
 gln ala glu leu ala lys pro gln gln ser val lys leu ser cys lys ala ser gln his 4411/1471 4441/1481 4471/1491
 AAA CAG AGG CCT GGA CAG GGT CTG GAA TGG ATT CGG TAC ATT AAT CCT AGC ACT ATT CCT AGT ATT ATT ATT ATT ATT ATT ATT ATT
 lys gln arg pro gln gln gln leu glu trp ile gln tyt ile asn leu ser ser gln tyt 4501/1501 4531/1511 4561/1521
 GCC ACA CTG ACT GCA GAC AAA TCC TCC AAC ACA CGC TAC ATT CAT CTG AGC ACC CTG ACA ATT GAG GAC TCT GCA GTC ATT TAC TGT GCA
 ala thr leu thr ala asp lys ser ser 4591/1531 4621/1541 4651/1551 peptide de
 AGG GCA GCT CAG GCT ACC ACC TTT GAC TAC TGG GGC CAA CGC ACC ACT CTC ACA GTC TCC TCA GGT GGA CGC CGT TCA CGC CGC CGT CGC
 arg ala ala gln ala thr phe asp tyt trp gln
 4681/1561 →: VL 4711/1571 4741/1581
 TCT GGC GGT GCG GGA TCC GAC ATT CTG ATG ATC CAG TCT CAC AAA TTC ATT TCC ACA TCA GTC CGA GAC AGG GTC AGC ATT ACC TGC AAG
 ser gln
 4771/1591 4801/1601 4831/1611
 GCC AGT CAG GAT GTG AGT ATT GCT GTC CGC TGG ATT CAA CAA AAA CGA GGG CAA TCT CCT AAA CTA CTG ATT TAC TGG GCA TCC ACC CGG
 ala ser gln esp val ser thr ala val ala trp tyt gln gln lys pro gln gln ser pro 4861/1621 4891/1631 4921/1641
 CAC ACT GCA GTC CCT GAT CGC TTC ACA CGC ACT GGA TCT GGG ACA GAT TAT ACT CTC ACC ATT AGC AGT GTC CAG CGT GCA GAC CTG GCA
 his the gln val pro asp arg phe the gln ser gln ser gln asp tyt thr leu thr 4951/1651 4981/1661 5011/1671
 CTC TGT TAC TGT CAG CAA CAT ATT AGC ACT CCT CGG ACC TTC GGT CGA CGC ACC AAC CGT CCT GGT GGA CGC GCA GGT GAA ATT AAA CGG CCT
 leu tyt tyt cys gln gln his tyt ser thr pro pro the gln gln gln gln lys leu glu ile lys arg ala pro gln gln gln gln gln
 ↓ 2gln 1Cys
 fin ScFv

CDR des régions variables VH et V

pACgp67-ScFv350 -> 1-phase Translation

Fig. 2C

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5041/1691 5071/1691 5161/1701
 aga tct gat cct TTC CTG GGA CCC GGC AAG AAC CAA AAA CTC ACT CTC TAC AAG GAA ATG CGT AAT GGT AAA CTC GAC ACG ATG AAG CTT
 arg ser esp pro phe leu gly pro sly lys asn sly lys leu thr leu phe lys sly ile arg asn val lys pro esp thr met lys leu
 5131/1711 5161/1721 5151/1731
 GTC GTC GGA TGG AAA GGA AAA GAG TTC TAC AGG GAA ACT TGG ACC CGC TTC ATG GAA GAC ATC CCC ATT GGT AAC GAC CAA GAA GTC
 val val sly trp lys gly lys sly the tyr arg sly the trp thr arg phe met sly asp ser phe pro ile val asn asp sly sly val
 5221/1741 5251/1751 5281/1761
 ATG GAT GTC TTC CTT GGT GTC AAC ATG CCT CCC ACT AGA CCC AAC CGT TGT TAC AAA TTC CGG CCT CAA CGC CCT CTG CGT TGC GAC CCT
 met esp val phe leu val val asn met arg pro thr arg pro asn arg cys tyr lys phe
 5211/1771 5341/1781 5371/1791
 GAC TAT GTC CCT CAT GAC GTG ATT AGG ATC GTC GAG CCT TCA TGG CTG CCT AGC AAC AAC GAG TAC CGC ATT ACC CTC CCT GTC GTC AAC GAG AAG
 asp tyr val pro his esp val ile arg ile val glu pro ser trp val sly ser asp asp sly glu tyr arg ile ser leu ala lys gly
 5401/1801 5431/1811 5451/1821
 GGC GGC TGC CCA ATA ATG AAC CCT CAC TCT GAG TAC ACC AAC TCG TTC GAA CAG TTC ATG GAT CGT GTC ATC TGG GAG AAC TTC TAC AAG
 gly sly cys pro ile met asn leu his ser glu tyr the asn ser phe glu sly phe ile esp arg val ile esp glu asn phe tyt lys
 5491/1831 5521/1841 5551/1851
 CCC ATC CCT TAC ATC GGT ACC GAC CCT GCT GAA GAG GAA ATT CTC CCT GAA CCT TCC CTC GTG TTC AAA GTC AAC GAG TTC GCA CCA
 pro ile val tyt ile sly the asp ser ala glu glu glu sly ile leu leu glu val ser leu val phe lys val lys sly phe ala pro
 5581/1861 5611/1871 5641/1881
 GAC GCA CCT CTG TTC ACT GGT CGG CGG TAT TAA AAC ACC ATA CAT TGT CAT TAG AAC ATT TAT TAA GCG CTA GAT TCT GTG CCT TGT TCA
 asp ala pro leu phe thr sly pro ala tyt OCH asn thr ile his cys tyt AMB tyt ile tyt OCH ala leu esp ser val arg cys OFA
 5571/1891 5701/1901 5731/1911
 TTT ACA GAC AAT TGT TGT AGG TAT TTT AAT ATT TCA TTA AAT TAA TAA TCT TTA CGG TGG TAC TGT AGA CGG AAA ATC AAA TCA TTT TCA
 the thr esp asn cys cys thr phe asp asn ser leu asp leu OCH ser leu gly tyt tyt val arg ala lys ile lys OFA phe ser
 5761/1921 5791/1931 5821/1941
 GCG TCT TTA TAT CTG AAT TAA AAT ATT AAA TCC TCA ATA GAT TGT TAA ATT AGG TTT CGA TTA GTT TCA AAC AAG GGT TGT TCT GAA
 ala ser leu tyt leu asn leu asn ile lys ser ser ile esp leu OCH asn arg phe arg leu val ser asn lys sly cys phe ser glu
 5851/1951 5881/1961 5911/1971
 CGG ATG GCT CGA CTA CCT ATT CGG TTT TCG CTC AAC ACA AAA CCT CCT AAC TGT TGT AGC AAC ATT CTA CCT TGG TCG AAT TTC GTC
 pro met ala gly leu ser asn sly phe ser leu asn ala thr lys leu ala lys ser cys
 5941/1981 5971/1991 6001/2001
 TGT GTT TTG TTT TGT AAT AAA CGT TCG AGC TCG TTC AAA ATA TAA TCA TGC CCT TTT GTC TTT 6031/2011
 CTT TCA TCA CTG TCG TTA GTG TAC ATT TGA TAA TCT TAA TCA TGG CCT AAC TTT CCT GTC TAA CCT GTC TCC TCA CCA CCC
 cys val leu phe cys asn lys sly ser thr ser phe lys ile leu cys ala phe val the leu ser ser leu ser leu val tyt esp OFA
 6061/2021 6091/2031
 CTC GAC GTC AAC ACG TTA AAT AAA CCT TGG ACA TAT TCA ACA TCG CGC GTG TAA CCT TTA leu asp val asn thr leu csn lys ala trp
 6121/2041 6151/2051 6181/2061
 TCG TCG TTA GAA GTT CCT TCC GAA GAC GAT TTT CGC ATA GCA CGA CCC CTC TAA ATT ser ser leu glu val ala ser glu val
 6211/2071 6241/2081 6271/2091
 GCA GTT GAG CTT TTT GGA ATT ATT CCT GAT TCC CGG CGT TTT TGG CGG CGT TTC ATT CTA val val glu leu phe gly ile ile ser esp
 6301/2101 6331/2111 6451/2151
 TTA GAA AGC GAC GGT GCA CGC CGT CGT AAC ATT TCA CAC CGC AAA CCT ACT ATT CCT CGC GGC leu glu ser esp sly ala sly sly asn
 6391/2131 6421/2141 6551/2171
 CGT GGA CGC GCA GGC CGC CGC CGA CGC CGC CGC CGT CGT CGC CGT GAT GCA GAC CGT CGT GCA CCT GTC TAA ATT CCT TCT TTA
 sly sly cly ala sly cly ala sly sly cly sly sly cly cly cly cly cly cly asp ala
 6481/2161 6571/2191 6651/2201
 CGC AAC'ACA' GTC CGC AAC TCA ACT ATT GTC CGT GTT CGC CCT GGT CCT GGT CCT GAT GCA GAC CGT CGT GAT GAT AAA CCT ACC ATC
 gly asn thr val gly thr ser thr ile val leu val ser gly ala val the gly leu the
 6661/2211 6691/2231 6651/2221
 CTA ATA CCT TCC AAC ATT TGT TGT CTG TCG TGT AAA GGT GCA CGC CGT CCT GTC GGT CCT CCT GTC ATT CCT GTC ATT CCT GTC ATT
 leu ile ala ser asn asn cys cys leu ser ser lys gly ala ala gly ser OPA gly ser val
 6661/2221 6691/2231 6721/2241
 ATT GAT'GCT' GGT GGT CGT CGT GCA CGC CGT GCA ATG TTA CGC ACC CGG GAA CCT
 ile asp gly sly sly sly ala gly met leu gly the gly glu gly gly gly
 6751/2251 6781/2251 6811/2271
 TTA GTT GTT TCG CGC ACC ATT CCT CGC ACC CGC GCA CGC CGT CGC CCT CCT TGT ACA ACC GAA leu val cys ser arg the ile val
 6841/2261 6871/2291 6901/2301
 CGT'GCG' AAC' TCA ATA TTA TAA TTG GAA TAC AAA TCG TAA AAA CCT
 sly sly asn ser ile leu OCH leu glu tyt lys ser OCH lys ser ala ile ser ile val
 6931/2311 6961/2321 6991/2331
 AAC AAC CGC TCA ATA TAA GCA ATT GTC TGG TAA AGA GAT TGT CTC AAG CTC CGC ACC CGT ATT CCT GTC ATT CCT CCT CCT CCT CCT
 asn asn arg ser met OCH ala ile val leu OCH arg esp cys leu lys leu arg the pro
 7021/2341 7051/2351 7081/2351
 TGG TAG TGG CGA GAC ACT TCG CTG TCG TCG ACC TAC ATG ATT CCT
 leu AMB tyt arg esp thr ser leu ser ser thr tyt met tyt ala leu leu ser lys the
 7111/2371 7141/2381 7171/2391
 ACA CCT CGT TTC ACC ACC ACT GTG TTG TCG TAA ATG CCT TGT TGT ACA ATT CCT
 thr ser leu phe ser thr thr val leu ser OCH met leu phe leu ile cys ala ser
 7201/2401 7231/2411 7261/2421
 GCA TCA ATT TIG TIG TCC CTA TTA TIG ATT AAA CCT
 ala ser ile leu leu phe leu leu asn lys OCH esp cys thr esp ser tyt leu arg
 7291/2431 7321/2441 7351/2451
 CAA AAC CGT TCA CAA ATT TAC GAA AAC TGC AAA AAC ACT CGG CCT ATT AAA ATA ATC AAC GGG CGC TTT CGC AAA ATA CCT ATT TAA
 sly the leu val sly phe tyt glu asn cys lys asn val lys the arg tyt lys ile ile
 7381/2461 7411/2471 7441/2481
 TGG CAC AAC CGC ACT AGC AAA CCT TAT TTG CAG AAA ACA ACT TCG CGG CGC AAC ATT CCT CCT GAC GAA ATA AAA CCT CAC CAG TAA ATC
 ser his lys pro thr ser lys leu tyt leu sly lys the ile ser ala his asn phe asp
 7471/2491 7501/2501 7531/2511
 AGC GAC CAC CCA ATT TTT ATA AAA ATT TAT CCT ATT CGT CCT TCC ACC AAC AAC CAA CCT ATT GTC ATG GAC TAC ATT GAC TGT CCT GAT
 ser esp his pro asn phe ile lys ile tyt phe asp his gly ser ile asn asn sly val
 7551/2521 7581/2531 7621/2541
 TTA TTT GAA ACA CTA GAA ATT AAA CGC GAG CCT TCG TAC CAA CCT GTT AGC ATT ATT AGA CAG CGT TGT GAA CGG CTC AAC GAT TAC
 leu phe sly the leu glu ile lys sly sly leu ser tyt sly leu val ser asn ile ile arg sly leu cys glu ala leu esp leu

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Fig 2D

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7651/2551 7681/2561 7711/2571
 CAC AAG CAC AAT TTC ATA CGC AAC GAC ATA AAA CTC GAA AAT GTC ATA TAT TTC GAA GCA
 his lys his asp phe ile his asn asp ile lys leu glu asn val leu tyt phe glu ala
 7741/2581 7771/2591 7801/2501
 TTG TGC AAA CGC GAA AAC TCA CTT AGC GTG CAC GAC GGC AGC TGC GAG TAC TTT AGT CGG
 leu cys lys his glu asn ser leu ser val his asp tyt ile leu glu tyt phe ser pro
 7831/2611 7861/2621 7891/2531
 TCG TTT GAC TGG TAC GCG GCG TGT TAA CAT ACA AGT TGC TAA CGG Gcg gtt cgt AAT GCT
 ser phe asp tyt phe ala ala cys OCH his the ser cys OCH pro ala val arg asn his
 7921/2541 7951/2651 7981/2661
 ATC CGC TCA CAA TTC CAC ACA ACA TAC GAG CCG GAA GCA TAA AGT GCA AAG CCT GGG GTG
 ile arg ser gln phe his the thr tyt glu pro glu ala OCH ser val lys pro gly val
 8011/2671 8041/2681 8071/2691
 CGT TGC CCT CGC TCC CGG CTT TCC AGT CGG GAA ACC TGT CCT CCC AGC TCC ATT AAT GAA
 arg cys ala his cys pro leu ser ser arg glu the cys arg ala ser cys ile asn glu
 8101/2701 8131/2711 8161/2721
 GTA TTG CGC CCT CTT CGG CTT CCT CGG TCA CGT ACT CGC TCC GGT CGG TGC GGC TGC
 val leu gly ala leu pro leu pro arg ser leu the arg cys ala arg ser phe gly cys
 8191/2731 8221/2741 8251/2751
 TAA TAC GGT TAT CCA CAG AAT CAG GGG ATA ACG CAG GAA AGA ACA TGT GAG CAA AAG GCG
 OCH tyt gly tyt pro gln est gln gly ile the gln glu arg the cys glu gln lys ala
 8281/2751 8311/2771
 CGT TCC TGG CGT TTT TCC ACA GGC TCC CGC CCT CGG TAC AGC AGC ATT ACA AAA AAT GAC CCT
 arg cys tyt arg phe ser ile gly ser ala pro leu the ser ile the lys ile asp ala
 8371/2791 8401/2801
 TAT AAA GAT ACC AGG CCT TTT CCC CTG GAA GCT CCC CGG TCC CCT CTC CGT CCC CGG TAA
 tyt lys asp the arg arg phe pro leu glu ala pro ser cys ala leu leu phe arg pro
 8461/2821 8491/2831 8521/2841
 TCC CCT CGG GAA GCG TGG CGC TTT CTC ATA GCT CAC CCT GAA GGT ATC TCA GTT CGG TGT
 ser leu arg glu ala tyt arg phe leu ile ala his ala val gly ile ser val arg cys
 8551/2851 8581/2861 8611/2871
 ACG AAC CCC CGG TTC AGC CGG ACC GCT CCT TAT CGG GAA ACT AAT GTC TGC AGT CCT CCA
 thr asn pro pro phe ser pro the ala ala pro tyt pro val the ile val leu ser pro
 8641/2861 8671/2891 8701/2901
 CAG CAG CCA CTG ATA ACA GGA TTA GCA GAG GGA GGT ATG TAG CGG GTG CTA CAG AST TCT
 gln gln pro leu val the gly leu ala glu arg gly met AMB ala val leu gln ser ser
 8731/2911 8761/2921 8791/2931
 GAA GCA CAG TAT TTG GTA TCT CGG CTC TGC TGA AGC CAG TTA CCT TTG GAA AAA GAG TTG
 glu gly gln tyt leu val ser ala leu cys OPA ser gln leu pro ser glu lys glu leu
 8821/2941 8851/2951 8881/2951
 CTG CTA CGG GTG GTT TTT TTG GCA AGC AGC AGA TTA CCT GCA GAA AAA AAG GAT CTC
 leu val ala val val phe leu phe ala ser ser arg leu arg ala glu lys lys asp leu
 8911/2971 8941/2981 8971/2991
 CTG AGC CCT AGT GGA AGC AAA ACT CAC GTT AAG GGA CCT TTG TCA TGA GAT TAT CAA AAA
 leu the leu ser gly the lys the his val lys gly phe tyt ser OPA asp tyt gln lys
 9001/3001 9031/3011 9061/3021
 AAT GAA GTT TTA AAT CAA TCT AAA GCA TAT ATG AGT AAA CCT GGT CTC ACA GTT ACC AAT
 asn glu val leu asn gln ser lys val tyt met ser lys leu gly leu the val the asn
 9051/3031 9121/3041 9151/3051
 TCT GTC TAT TTC GTT CAT CCA TAC TAC CCT GAC TCC TCG TGT AGA TAA CTA CGA TAC
 ser val tyt phe val his pro AMB leu tyt asp ser pro ser cys arg OCH leu arg tyt
 9181/3061 9211/3071 9241/3081
 CAA TGA TAC CGG GAG ACC CAC GCT CAC CGG CCT CAG ATT TAT CAG CAA TAA ACT ACC CAG
 gln OPA tyt arg glu the his ala his arg leu gln ile tyt gln gln OCH the ser gln
 9271/3091 9301/3101 9331/3111
 CAA CCT CCT CCT CCA TCC AGT CTA TTA ATT CCT GGT CGG AAG CTA GAG TAA GCA GTT
 gln leu tyt pro pro ser ser leu leu ile val ala gly lys leu gln OCH val val
 9361/3121 9391/3131 9421/3141
 CCA TAC CTA CAG GCA CGG TGG TGT CAC CCT CGT CGT TGG GAA TGG CCT CAT TCA GCT CGG
 pro leu leu gln ala ser tyt cys his ala arg arg leu val tyt leu his ser ala pro
 9451/3151 9481/3151 9511/3171
 CCC TGT TGT GCA AAA AAG CGG TTA GCT CCT TCG GTC CTC CGA TCC TGC TCA GAA GCA
 pro pro cys cys ala lys lys arg leu ala pro ser val leu arg ser leu ser glu val
 9541/3161 9571/3191 9601/3201
 TCG CAG CAC TGC ACA ATT CTC TCA CTG TCA TGC CAT CCT CGG TAA GAT GCT TTT CTC TGA CTG
 tyt gln his cys ile ile leu leu leu ser cys his pro OCH asp ala phe leu OPA leu
 9631/3211 9661/3221 9691/3231
 AGT GTC TGC GGC GAC CGA GGT CCT CCT CGG CCG CGT CAA TAC CGG ATA ATA CCT CGG CAC
 ser val cys gly asp arg val ala leu ala arg arg gln tyt gly ile ile pro arg his
 9721/3241 9751/3251 9781/3251
 GAA AAC CCT CCT CGG CGC GAA AAC TCT CAA CGA CCT TCT TAC CGG TGT TGA GAT CGA GGT CGA
 glu asn val leu arg gly glu asn ser gln gly ser tyt arg cys OPA asp pro val arg
 9811/3271 9841/3281 9871/3291
 CAC CAT CCT TTA CCT TCA CCA CGG TTT CGT GGT GAG CAA AAA CGA GAA CGC AAA ATT CGG
 gln his leu leu leu ser pro ala phe leu gly glu gln lys gln glu gly lys net pro
 9901/3301 9931/3311 9961/3321
 GTT GAA TAC TCA TAC TCT TTT TTC ATA ATT ATT GAA GCA TTT ATA AGG GTT ATT GTC
 val gln tyt ser tyt ser ser phe the ame ile ile glu ala phe ile arg val ile val
 9991/3331 10021/3341 10051/3351
 AGA AAA ATA AAC AAA TAG CGG TTC CGC GCA CAT TTC CCT CGG CGA AGG TGC CAT CGT ACC TCT
 arg lys ile asn lys AMB gly phe arg ala his phe pro gln lys cys his leu the ser
 10021/3361 10111/3371 10141/3381
 ACA AAA AEA GGC GTC TCA CGA CGC CCT TTC GTC CGC CCT TCC CGT AGG AGG CGT AAA
 ile lys ile gly val ser arg gly pro phe val ser arg val ser val met the val lys
 10171/3391 10201/3401 10231/3411
 TCA CGG CCT GTC TGT AAG CGG ATG CGG GCA GAC AGG CGC CCT CGG CGT CGC TTA ACT
 ser gln leu val cys lys arg net pro gly ala esp lys pro val arg ala arg gln arg
 val leu ala gly val gly ala gly leu the

pACgp67-ScFv350 -> 1-phase Translation

Fig 2E

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10281/3421 10291/3431 10321/3441
 ATG CGG CAT CAG AGC AGA TTG TAC TGA GAG TGC ACC ATG TGC GGT GTG AAA TAC CGC ACA GAT CGG TAA GGA GAA AAT ACC GCA TGA GGC
 met arg his gln ser arg leu tyt OFA s:lu cys thr ile cys gly val lys tyt arg thr asp ala OCH sly s:lu asn thr ala ser gly
 10351/3451 10381/3461 10411/3471
 GCC ATT CGC CAT TCA GGC TGC GCA ACT GTT GGG AAG CGC GAT CGG TGC CGC CCT CTT CGC TAT TAC CGC AGC TGG CGA AAG CGG GAT GTG
 ala ile arg his ser gly cys ala thr val gly lys gly asp arg cys gly pro leu arg tyt tyt ala ser trp arg lys gly asp val
 10441/3481 10471/3491 10501/3501
 CTG CAA GGC GAT TAA GTF GGG TAA CGC CAG GGT TTT CCC AGT CAC GAC GTT CTA AAA CGA CGG CCA GTG CC
 leu gln gly asp CCA val gly gln gly phe pro ser his asp val val lys arg arg pro val

Figure 3 : clonetherap.99B3**VH sequence :**

GAGGTGAAGCTTCTCCAGTCTGGAGGTGGCCTGGTGCAGCCTGGAGGGATCCCTGA
AACTCTCCTGTGCAGCCTCAGGAATCGATTTAGTAGATACTGGATGAGTTGGGT
TCGGCGGGCTCCAGGGAAAGGACTAGAAATGGATTGGAGAAATTAATCCAGATAG
CAGTACAATAAACTATGCACCATCTCTAAAGGATAAAATTCACTCATCTCCAGAGAC
AACGCCAAAAATACGCTGTACCTGCAAATGAGCAAAGTGAGATCTGAGGACACA
GCCCTTATTACTGTGCAAGAGGACTGGGACAGAACTTGACTACTGGGGCCAAG
GCACCACTCTCACAGTCTCCTCA

VL sequence :

GATATTGTGATGACGCAGGCTGCATTCTCCAATCCAGTCACTCTTGGAACATCAG
CTTCCATCTCCTGCAGGTCTAGTAAGAGTCTCCTACATAGTAATGGCATCACTTAT
TTGTATTGGTATCTGCAGAACGCCAGGCCAGTCTCCTCAGCTCCTGATTTATCAGAT
GTCCAACCTTGCTTCAGGAGTCCCAGACAGGTTCACTAGCAGTGGGTCAAGGAAC
GATTCACACTGAGAATCAGCAGAGTGGAGGCTGAGGATGTGGGTGTTATTACT
GTGCTCAAAATCTAGAACTTCCGTGGACGTTGGTGGAGGCACCAAGCTGGAAAT
CAAA

Figure 4 : clonetherap.88E10**VH sequence :**

GAGGTGAAGCTGGTGGAGTCTGGAGGAGGCTTGGTACAGCCTGGGGTTCTCTG
AGTCTCTCCTGTGCAGCTTCTGGATTCACCTCACTGATTACTCCATGAACCTGGGT
CCGCCAGCCTCCAGGAAGACACTTGAGTGGTTGGCTTTATTAGAAACAAAGCT
AATGGTTACACAGCAGAGTACAGTGCATCTGTGAAGGGTCGGTTCTCCATCTCCA
GAGATAATTCCCAAAGCATTCTATCTTCAAATGAATGCCCTGAGAGCTGAGGA
CAGTGCCACTTATTACTGTGCAAGGGATGGTATGCTATGGACTACTGGGGTCAA
GGAACCTCAGTCACCGTCTCCTCA

VL sequence :

Figure 5 : clonetherap.152C3**VH sequence :**

GAGGTTCTGCTGCAGCAGTCTGTGGCAGAGCTTGTGAGGCCAGGGGCCTCAGTCA
AGTTGTCTGCATAGTTCTGACTTCAACATTAAACACACCTATATGCACTGGGTG
AAACAGAGGCCTGATCAGGGCCTGGAGTGGATTGGAAGGATTGATCCTGCGAAT
GGTAAAACATATATGCCCCACGTTCCAGGGCAAGGCCACTATAACTGCGGACA
CATCCTCCAACACAGCCTACCTGCATTTCAAGCAGCCTGACATCTGAGGACGCTGC
CATCTATTACTGTGCTAGAGCTGGGCTGGCTACTTTGACTACTGGGGCCAAGGC
ACCACTCTCACAGTCTCCTCA

VL sequence :

GACATCTTGTGACTCAGTCTCCAGCCATCCTGTCTGTGAGTCCAGGAGAAAGAG
TCAGTTCTCCTGCAGGGCCAGTCAGAACATTGGCACAAGTATTACTGGTATCA
GCAAAGAACAAATGGTTCTCAAGGCTTCTCATAAAGTATGTTCTGAGTCTATC
TCTGGGATCCCTCCAGGTTAGTGGCAGTGGATCAGGGACAGAGTTACTCTTA
GCATCAACAGTGTGGAGTCTGAAGATATTGCAGATTATTACTGTCAACAAAGTCA
TAGTTGGCCGCTCACGTTCGTGTGGACCAAGCTGGAGCTGAAA

Fig. 6

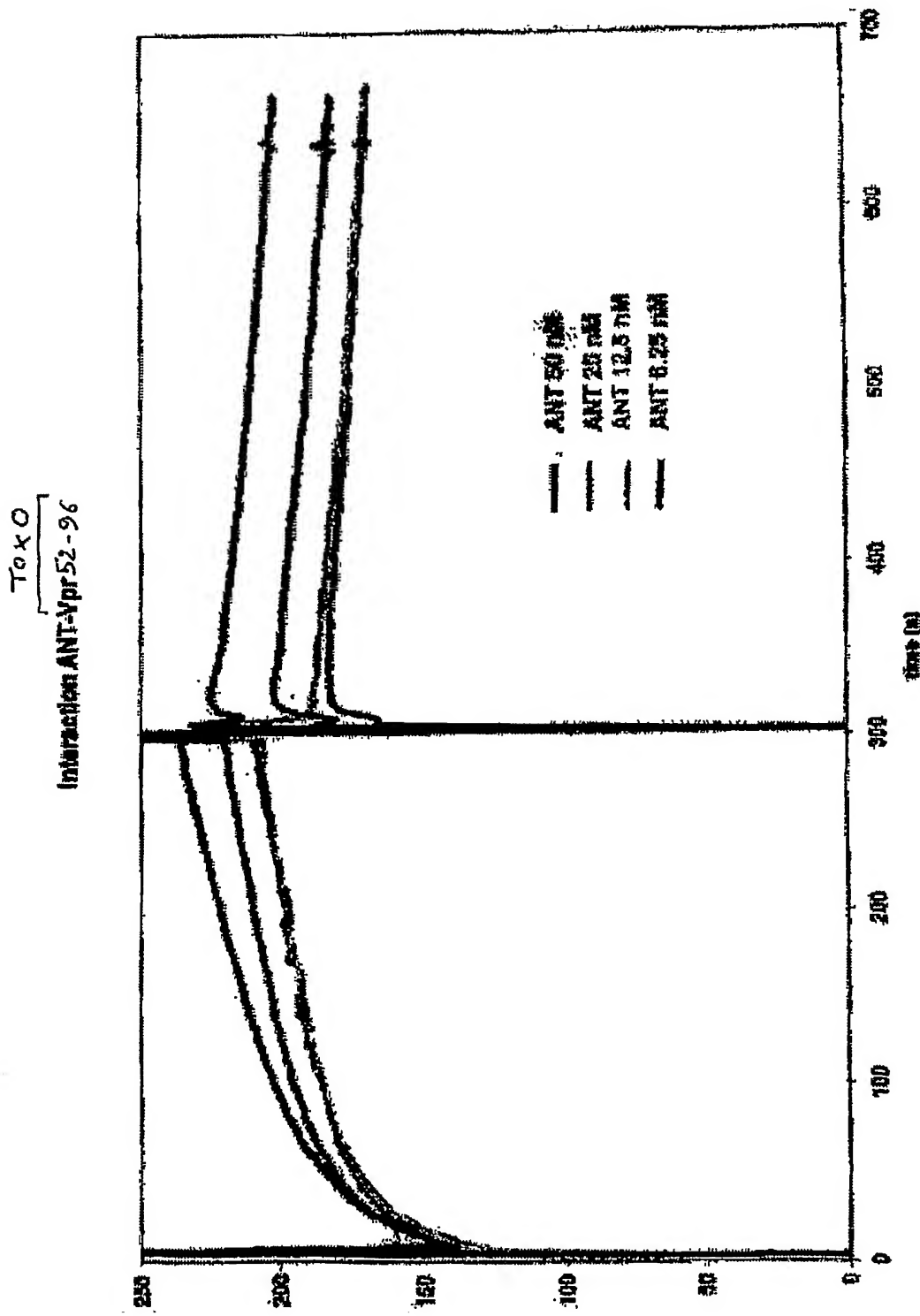


Fig. 7

Untersuchung AHT-Toxi

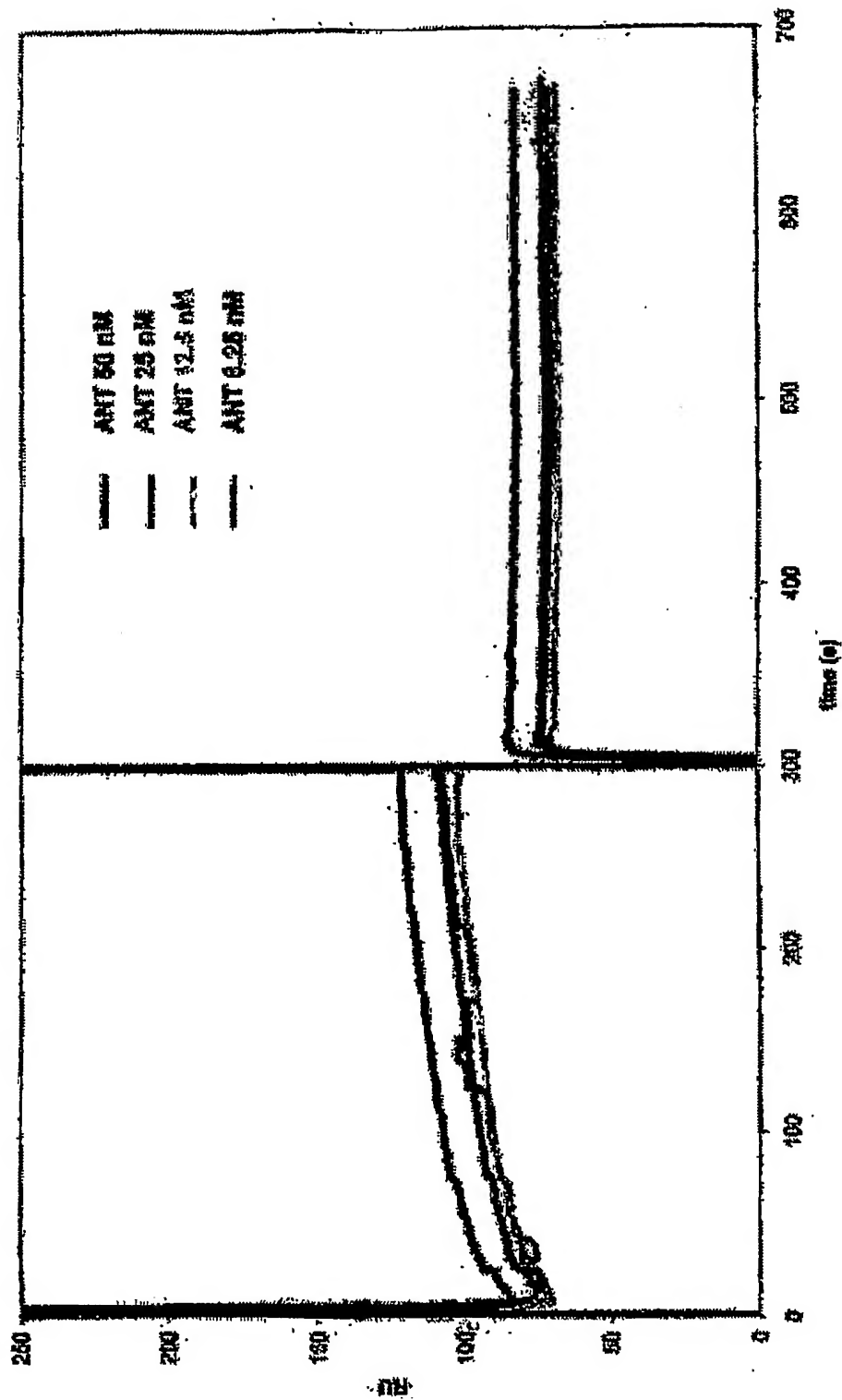


Fig. 8

$\overbrace{\text{Tox 0}}^{\text{Interaction VDAC-Yip52-96}}$

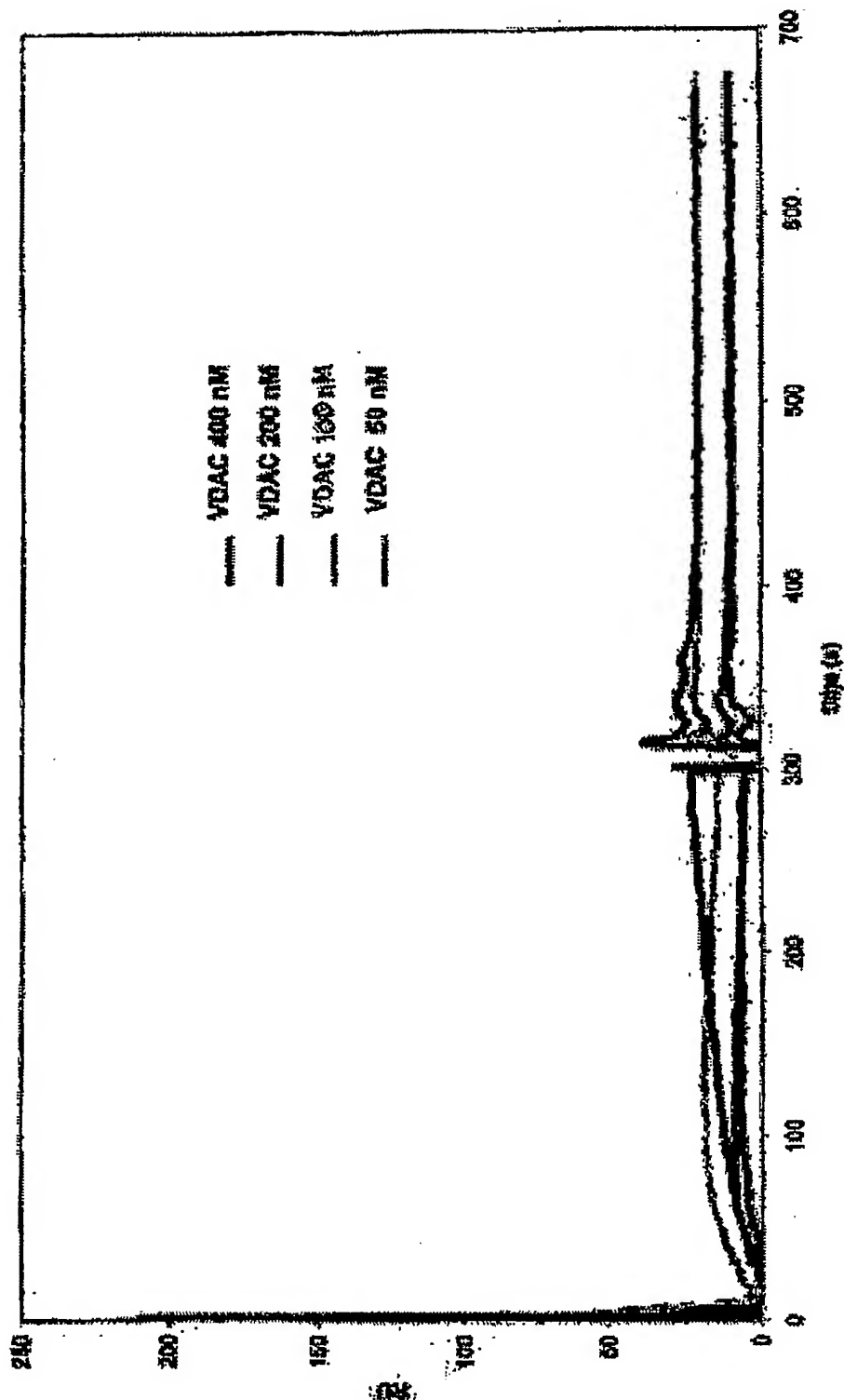


Fig. 9

Interaction VDAC-Tort

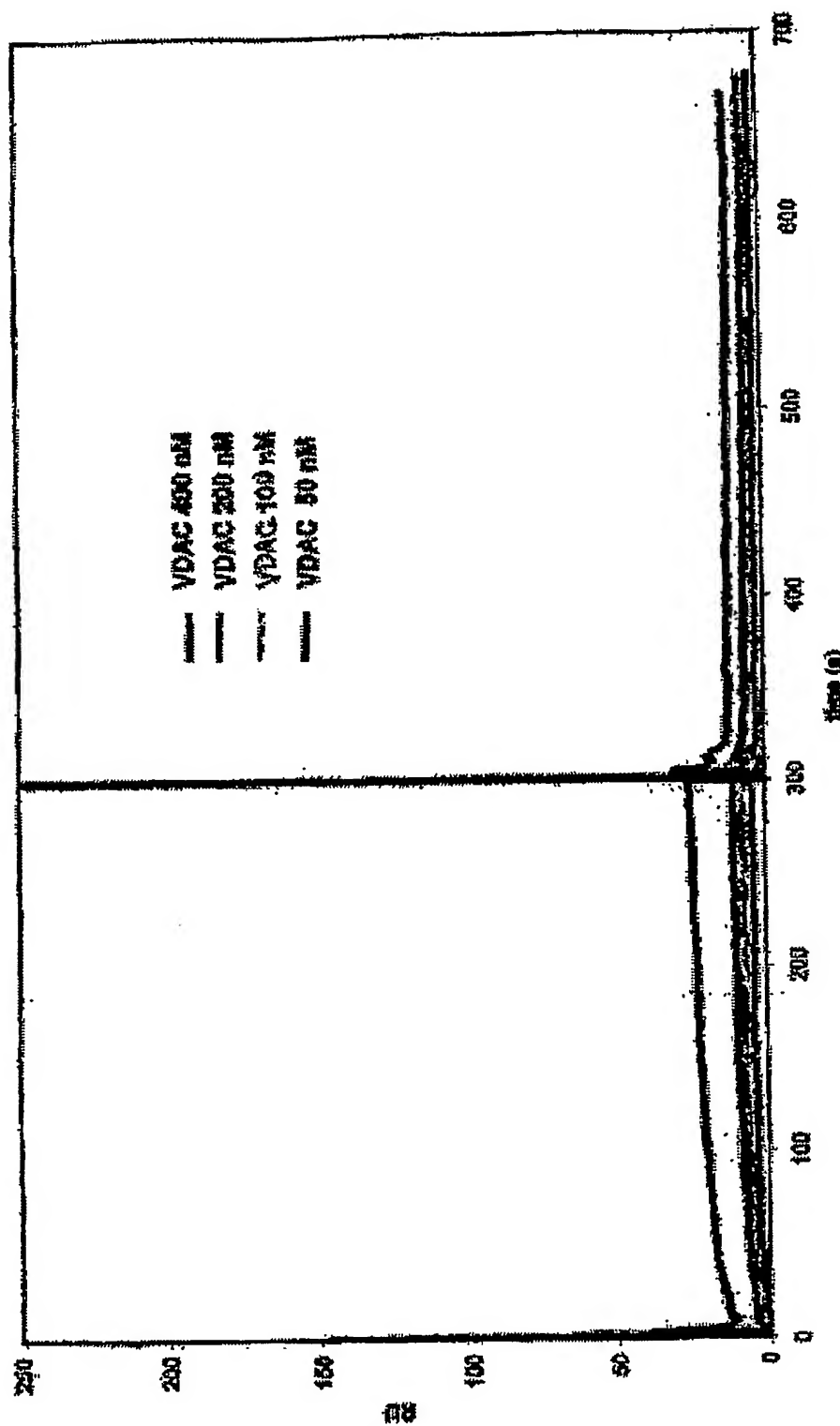


Fig. 10

Interaction RHT 80 rpm - prepared

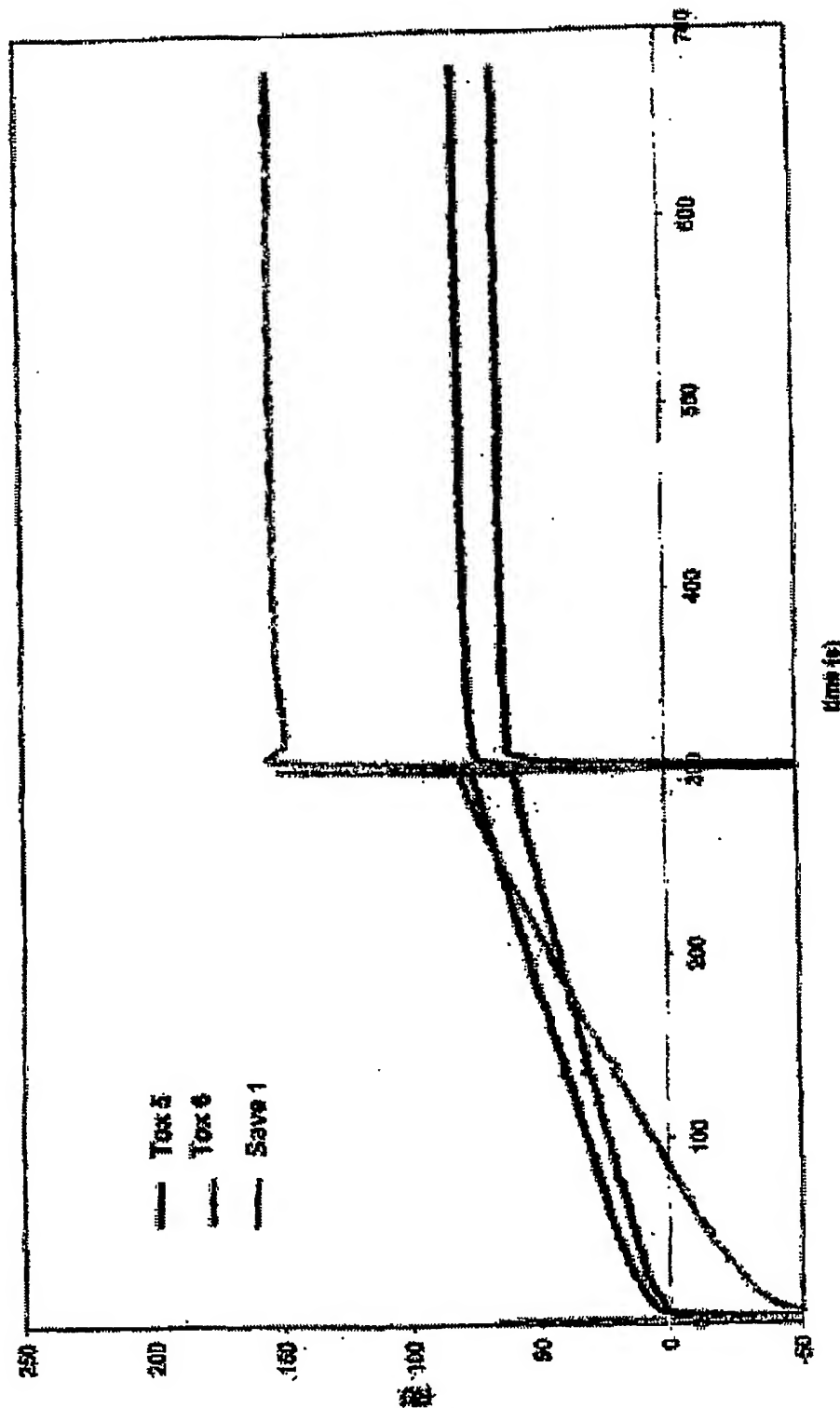
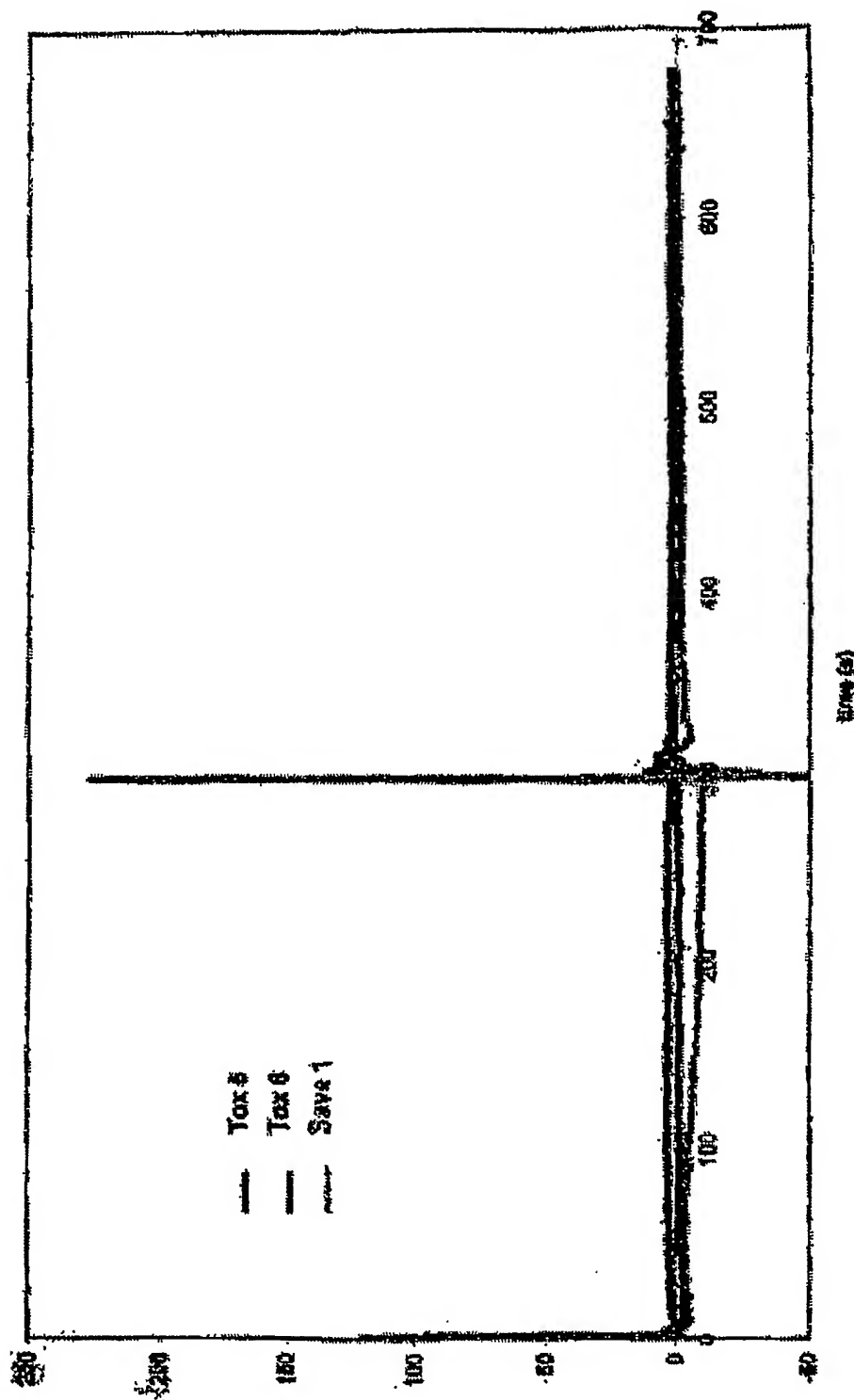


Fig. 41

Interaction VDAC 50 nM + peptide



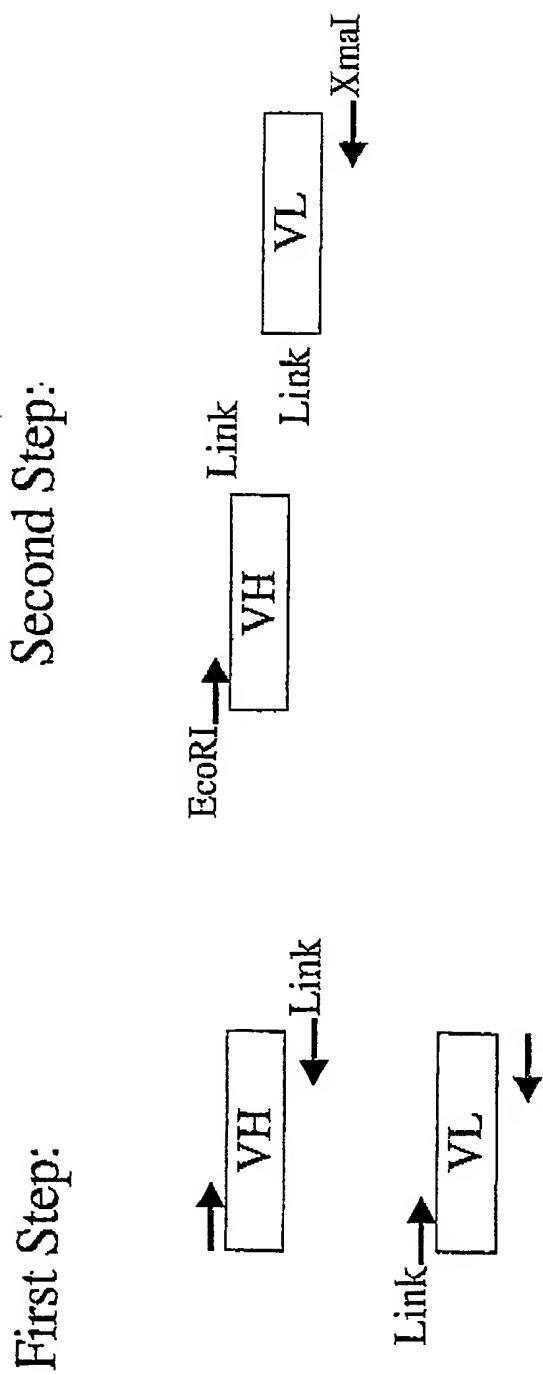


Figure 12: Obtention of the VH/VL chimeric DNA

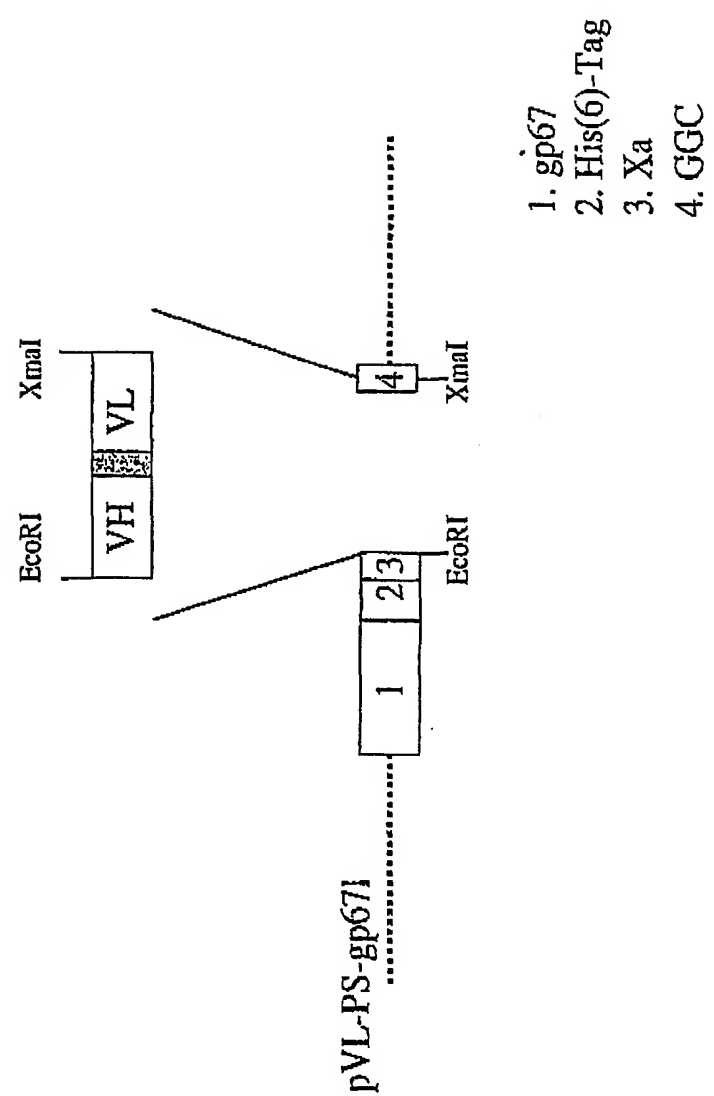


Figure 13: Map of the ScFv transfert vector